PRM PTO-1390	S DEPARTMENT OF COMMERCE PATENT AND TRADEMA				
EV 10-94)  TRANSMITTAL LETTE	ARK OFFICE ATTORNEY'S DOCKET NUMBER 11613.33USWO				
DESIGNATED/ELEC CONCERNING A FIL					
		Us APPLICATION NO (If known, see 37 C F R 1 5) Unkhowh / O I O O O			
TERNATIONAL APPLICATION NO.	INTERNATIONAL FILING DATE	PRIORITY DATE CLAIMED			
CT/US00/14489	25 May 2000	18 June 1999			
TLE OF INVENTION					
DK4 BINDING PEPTIDE					
PPLICANT(S) FOR DO/EO/US					
ATERSON et al.					
oplicant herewith submits to the United States	s Designated/Elected Office (DO/EO/US) the	following items and other information:			
[X] This express request to begin national examination until the expiration of the [X] A proper Demand for International Propertion of the International Application and [X] is transmitted herewith (requestive by the International for Internationa	UENT submission of items concerning a filing examination procedures (35 U.S.C. 371(f)) at applicable time limit set in 35 U.S.C. 371(b) eliminary Examination was made by the 19th on as filed (35 U.S.C. 371(c)(2)) ired only if not transmitted by the International nuternational Bureau. Application was filed in the United States Received Application into English (35 U.S.C. 371(c)(2)) mational Application under PCT Article 19 (35 (required only if not transmitted by the International Bureau. Wever, the time limit for making such amendment of the claims under PCT Article 19 (35 U.S.C. on the claims under PCT Article 1	any time rather than delay and PCT Articles 22 and 39(I). month from the earliest claimed priority date.  Il Bureau).  Ving Office (RO/US)  3.  5. U.S.C. 371(c)(3))  ational Bureau).  Dents has NOT expired.  371(c)(3)).			
(35 U.S.C. 371(c)(5)).					
ns 11. to 16. below concern document(s) or information included: [ ] An Information Disclosure Statement under 37 CFR 1.97 and 1.98.					
[ ] An assignment document for reco	rding. A separate cover sheet in compliance w	ith 37 CFR 3.28 and 3.31 is included.			
A FIRST preliminary amendment. A SECOND of SUBSEQUENT preliminary amendment.					
[ ] A substitute specification.					
] A change of power of attorney and/or address letter.					
[X] Other items or information: Form PCT/IPEA/416					
		i i			

. APPLICATION NO. (If kno	own, see 37 C F R 1 5)	INTERNATIONAL APPLICATION	531 R	ec'd Pari	18 DEC 2001
Unknown 1	0/018964	PCT/US00/14489		11613.33USWO	ER C
17. [X] The following				CALCULATIONS	PTO USE ONLY
BASIC NATIONAL F	FEE (37 CFR 1.492(a) (1)-(	(5)):			
Search Report has	been prepared by the EPO	ог ЈРО	\$890.00		
International preliminary examination fee paid to USPTO (37 CFR 1.492(a)(1))\$710.00					
No international probut international se	eliminary examination fee pearch fee paid to USPTO (3	aid to USPTO (37 CFR 1.47 CFR 1.445(a)(2))	82) <b>\$740.00</b>	,	
Neither international international search	al preliminary examination in the (37 CFR 1.445(a)(3)) p	fee (37 CFR 1.482) nor paid to USPTO	\$1040.00		
International prelim and all claims satis	ninary examination fee paid fied provisions of PCT Arti	to USPTO (37 CFR 1.482)	\$100.00		
ENTER APPROPRIATE BASIC FEE AMOUNT =				\$ 890.00	
urcharge of \$130.00 fo nonths from the earliest	r furnishing the oath or decl claimed priority date (37 C	aration later than [] 20 []. FR 1.492(e)).	30	\$	
CLAIMS	NUMBER FILED	NUMBER EXTRA	RATE		
otal claims	9 -20=		X \$18.00	\$	
dependent claims	6 -3 =	3	X \$84.00	\$ 252.00	
IULTIPLE DEPENDE	NT CLAIM(S) (if applicabl	e)	+ \$260.00	\$	
	TOTAL	OF ABOVE CALCU		\$1142.00	
	ng by small entity, if applica				
ırsuant 6 37 CFR 1.27	7	ore. Small cherry status is	Ciamico	\$	
		ST	JBTOTAL =	\$1142.00	
occessing fee of \$130.00 onths from the earliest	0 for furnishing the English claimed priority date (37 CF	translation later than [ 1 20		\$	
#500 0 #500 0 #500 0 #500		TOTAL NATIO	NAL FEE =	\$1142.00	
the for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be comparated by an appropriate cover sheet (37 CFR 3.28, 3.31). \$40.00 per property				\$	
		TOTAL FEES EN	CLOSED =	\$1142.00	
				Amount to be:	
			į	refunded	\$
				charged	\$
[X] Check(s) in the a	amount of \$1142.00 to cove	r the above fees is enclosed	•		
[ ] Please charge my Deposit Account No in the amount of \$ to cover the above fees.  A duplicate copy of this sheet is enclosed.					
[X] The Commission overpayment to l	ner is hereby authorized to c Deposit Account No. <u>13-27</u> 2	harge any additional fees w 25.	hich may be requ	ired, or credit any	
OTE: Where an appro 37(a) or (b)) must be f	opriate time limit under 37 filed and granted to restor	CFR 1.494 or 1.495 has rethe application to pending	ot been met, a p	etition to revive (37 CFR	
D ALL CORRESPONDENCE TO		•		1 /2	
Ielissa J. Pytel					I(X) = I
IERCHANT & GOUL P.O. Box 2903	D ·		SIGN	IATURE:	MINI
Ainneapolis, MN 5540	2-0903		NAM	IE: Melissa J. Pytel	)
				•	
			REG	ISTRATION NUMBER:	41,512

S/N 10/018964

PATENT

# IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant:

PATERSON et al.

Docket No.:

11613.33USWO

Serial No.:

10/018964

Filed:

18 December 2001

Int'l Appln No.:

PCT/US00/14489

Int'l Filing Date:

25 May 2000

Title:

CDK4 BINDING PEPTIDE

**CERTIFICATE UNDER 37 CFR 1.10** 

'Express Mail' mailing label number: EV036308473US

Date of Deposit: 11 April 2002

I hereby certify that this correspondence is being deposited with the United States Postal Service 'Express Mail Post Office To Addressee' service under 37 CFR 1.10 on the date indicated above and is addressed to the Assistant Commissioner for Patents, Washington, D.C. 20231.

Name: Chris Stordahl

# SUBMISSION OF COMPUTER READABLE SEQUENCE LISTING

**BOX SEQUENCE** 

**Assistant Commissioner for Patents** Washington, D.C. 20231

Dear Sir:

HODIMION LOUIS

This paper is being filed in accordance with the requirements of 37 C.F.R. § 1.821 - § 1.825. Applicants enclose herewith a computer readable form of the Sequence Listing. In addition, a paper copy of the Sequence Listing is also provided. Please insert the paper copy into the file after the Executed Oath or Declaration, as required by 37 C.F.R. § 1.77(a)(15). Applicants hereby state that the computer readable form and the paper copy of the Sequence Listing enclosed do not include new matter.

Respectfully submitted,

MERCHANT & GOULD P.C.

P.O. Box 2903

Minneapolis, Minnesota 55402-0903

(612) 332-5300

Dated: 11 April 2002

Reg. No. 33,112

JJG/kjr



10

15

20

25

30

35

# 1 531 Rec'd PCT/PT 18 DEC 2001 CDK4 BINDING PEPTIDE

# **BACKGROUND OF THE INVENTION**

The division cycle of eukaryotic cells is regulated by a family of protein kinases known as cyclin—dependent kinases (CDKs). The sequential activation of individual members of this family and their consequent phosphorylation of critical substrates promotes orderly progression through the cell cycle. CDK activities are regulated by cyclin binding, by both positive and negative regulatory phosphorylations and by polypeptide CDK inhibitors. Cellular differentiation is accompanied by the down—regulation of CDK activity, which occurs through at least two mechanisms: down—regulation of cyclin expression which is required for CDK activity; and induction of CDK inhibitor expression. CDK4 is a major catalytic subunit of mammalian D—type cyclins, which act during the G<sub>1</sub> phase of the cell cycle to enforce the decision of cells to enter the S phase. The deregulation of eukaryotic CDKs is correlated with many types of cancers.

Three D-type cyclins (D1, D2 and D3) are differentially expressed in proliferating cells in response to various growth factor mediated signals. The D-type cyclins interact combinatorially with CDKs 2, 4, 5 and 6 to form active holoenzymes that facilitate progression through the G<sub>1</sub> phase of the cell cycle into S phase. Expression of the D-type cyclins depends on mitogenic stimulation, regardless of the position of the cell in the cycle. Growth factor withdrawal leads to rapid cyclin D destruction, with an associated loss of CDK activity.

A classical example of terminal differentiation is that of skeletal muscle cells. Skeletal muscle differentiation entails the coordination of muscle specific gene expression and terminal withdrawal from the cell cycle. Muscle differentiation is intimately coupled to the cell cycle, such that muscle—specific transcription is initiated only when myoblasts are growth arrested in the  $G_1/G_0$  phase.

MyoD is a basic helix-loop-helix (bHLH) protein which plays an important role in the differentiation of skeletal muscle by inducing muscle structural gene expression and cell cycle withdrawal. In particular, MyoD is one of several transcriptional activators of muscle specific gene expression. The mechanisms by which MyoD induces myogenesis involve both the activation of muscle—specific gene expression and withdrawal from the cell cycle. Although proliferating myoblasts express MyoD throughout the cell cycle, the functions of MyoD are repressed in proliferating myoblasts.

10

15

20

25

30

Tumor suppressor retinoblastoma protein (RB) suppresses cell proliferation and thus plays an important role in the production and maintenance of the terminally differentiated phenotype of muscle cells. RB also appears to participate in control of entry into the S phase of the cell cycle. Inactivation of RB in terminally differentiated cells allows the cells to reenter the cell cycle. Correspondingly, loss of a functional RB is key step in the development of many human tumors.

The activity of RB is modulated by a phosphorylation/dephosphorylation mechanism during cell proliferation and differentiation. In resting or differentiated cells, RB protein is present in its dephosphorylated from. Underphosphorylated RB inhibits growth promoting transcription factors in the E2F / DP family. In rapidly proliferating cells, RB protein is highly phosphorylated. Maximal phosphorylation is associated with S phase of the cell cycle. Cyclins and CDKs contribute to the sequential phosphorylation of RB to inactivate its growth suppressive function.

#### **SUMMARY OF THE INVENTION**

The inventors have identified a peptide (hereinafter referred to as a "CDK4 binding peptide") that is capable of specifically binding a cyclin dependent kinase (CDK4) to inhibit CDK4 activity and cell growth. In one embodiment, the invention provides an isolated nucleic acid sequence comprising DNA encoding the CDK4 binding peptide or a nucleic acid sequence that is complementary to the isolated nucleic acid sequence and remains stably bound to it under at least moderate, and optionally, under high stringency conditions.

In another embodiment, the invention provides an isolated CDK4 binding peptide [SEQ. ID NO: 1]. The invention also includes variants of the CDK4 binding peptide which comprise polypeptides which have at least about 80% amino acid sequence identity with the amino acid sequence of the CDK4 binding peptide.

In another embodiment, the invention provides chimeric molecules comprising a CDK4 binding peptide fused to a heterologous peptide or amino acid sequence, preferably a nuclear localization signal.

Therapeutic and diagnostic methods are also provided.

#### BRIEF DESCRIPTION OF THE FIGURES

Figure 1 is a photograph of a gel showing that MyoD binding to CDK4 does not depend upon cyclin Dl.

10

15

20

25

30

Figure 2 is a graph showing the <u>in vivo</u> interaction between MyoD and CDK4 in the mammalian two-hybrid system.

**Figure 3** is a gel showing the interaction of MyoD and CDK4 in dividing C2Cl2 myoblasts.

**Figure 4** is a photograph of a gel showing that CDK4 specifically inhibits MyoD–E12 DNA–binding in the absence of cyclin Dl.

Figure 5 is a graph showing that CDK4 differentially inhibits MyoD and myogenin activation of the MCK CAT reporter plasmid in 10T1/2 cells grown in differentiation medium (DM).

**Figure 6** is a graph showing forced nuclear expression of CDK4 inhibits myogenic conversion of 10T1/2 cells in differentiation medium (DM) in the absence of cyclin Dl.

**Figure 7** is a photograph of a gel showing that MyoD and RB are not phosphorylated comparably by CDK4 in vitro.

**Figure 8a** is a graph showing that gal 4-RB and vp16-E2F interact to activate luciferase expression 10-15 fold routinely, whereas vp16-MyoD and vp16-myogenin show no significant activation above background with gal4-RB.

Figure 8b is a graph showing that both vp16-MyoD and vp16-myogenin interact strongly with gal4-E12.

Figure 9 is a schematic characterization of the CDK-binding domain in MyoD(CMD1) and regions used to study MyoD-CDK4 interactions.

Figure 10a is a photograph of a gel showing that CDK4 binds to the carboxy terminus of MyoD (lane 5) but does not bind to the other vertebrate myogenic factors; GST myogenin shown as an example (lane 3).

Figure 10b is a photograph of a gel showing that a fifteen amino acid domain between amino acid residues 189–203 that is required for CDK4 binding (lanes 4 and 5).

Figure 10c is a photograph of a gel showing that the fifteen amino acid CDK4-binding domain in MyoD (aa 189–203) binds CDK4 as efficiently as full-sized MyoD when joined to GST (lanes 2 and 3).

Figure 11a is a photograph of a gel showing that extracts from Sf9 cells coinfected with wild type virus (lane 1), a cyclin D1 encoding virus (lane 2) or a CDK4 encoding virus (lane 3) give background levels of kinase activity on serine 780 (antibody to phosphoserine 780) whereas coinfections with baculovirus stock encoding CDK4 and cyclin D1 produce active CDK4 (lane 4) that specifically phosphorylates RB (GST RB (aa 767–928)) on serine 780. This phosphorylation is

35

10

15

20

25

30

inhibited to background levels by the CDK specific kinase inhibitor p16 (lane 5, added as GST-p16).

Figure 11b is a photograph of a gel showing that full sized MyoD(CMD1) (second row), the C-terminus of MyoD(CMD1) (third row) and the fifteen amino acid CDK4 binding domain of MyoD fused to GST (fourth row) inhibit the CDK4-dependent phosphorylation of RB(GST-RB) in vitro whereas myogenin (first row) has no inhibitory affect).

Figure 12 is a table showing the growth inhibitory effects of the various MyoD(CMD1) subdomains joined to either the pCDNA3 or GST-pCEFL vectors. The number of BrdU positive cells per 500 lacZ positive cells, normalized to vector and averaged for six independent assays is shown. In the top panel, constructs were expressed with the pCDNA3 vector. In the lower panel constructs were fused to eukaryotic expression vector, GST-pCEFL, and contained a nuclear localization signal.

Figure 13 is a photograph of a gel showing the inhibition of RB phosphorylation in C2C12 myoblasts by MyoD(CMD1) and the C-terminus containing the fifteen amino acid CDK4 binding peptide.

Figure 14 is a sequence comparison of the fifteen amino acid CDK4 binding peptide in chicken MyoD(CMD1) versus human, rat, mouse, pig, xenopus and zebra fish. The consensus is given between chicken and mammals.

## **DETAILED DESCRIPTION**

#### **Definitions**

The term "CDK4 binding peptide", when used herein, encompasses both the native sequence shown and variants thereof (which are further defined herein). The CDK4 binding peptide may be isolated from a variety of sources, such as vertebrate sources, including human tissue types, or from another source, or prepared by recombinant or synthetic methods.

A "native" CDK4 binding peptide sequence comprises a polypeptide having the same amino acid sequence as a CDK4 binding peptide derived from nature. Native sequences can be isolated from nature or can be produced by recombinant or synthetic means.

A "variant" of the CDK4 binding peptide includes peptide sequences having at least about 80% amino acid sequence identity with the CDK4 binding peptide

shown in [SEQ. ID NO: 1], more preferably about 90%, even more preferably about 95% sequence identity.

"Percent (%) amino acid sequence identity" with respect to the CDK4 binding peptide sequence is defined as the percentage of amino acid residues in a candidate sequence that are identical to the amino acid residues in the CDK4 binding peptide sequence, after aligning the sequences and introducing gaps, if necessary to introduce maximum percent sequence identity, and not considering any conservative substitutions as part of the sequence identity. Alignment for purposes of determining percent amino acid sequence identity can be achieved in various ways that are within the skill in the art, for instance, using publicly available computer software such as BLAST, ALIGN or Megalign (DNASTAR) software. Those skilled in the art can determine appropriate parameters for measuring alignment, including any algorithms needed to achieve maximal alignment over the full length of the sequences being compared.

"Percent (%) nucleic acid sequence identity" with respect to the nucleic acid sequences encoding the CDK4 binding peptide is defined as the percentage of nucleotides in a candidate sequence that are identical to the nucleotides in the nucleic acid sequence encoding the CDK4 binding peptide after aligning the sequences and introducing gaps, if necessary, to achieve the maximum percent sequence identity. Alignment for the purpose of determining percent nucleic acid sequence identity can be achieved in various ways that are within the skill in the art, for instance, using publicly available computer software such as BLAST, ALIGN, or Megalign (DNASTAR) software. Those skilled in the art can determine appropriate parameters for measuring alignment, including any algorithms needed to achieve maximal alignment over the full length of the sequences being compared.

"Isolated" when used to describe the peptides disclosed herein, means a polypeptide that has been identified and separated and/or recovered from a component of its natural environment. Contaminant components of its natural environment are materials that would typically interfere with diagnostic or therapeutic use of the polypeptide, and may include enzymes, hormones, and other proteinaceous or non-proteinaceous solutes. In preferred embodiments, the polypeptide is purified (1) to a degree sufficient to obtain at least 15 residues of N-terminal or internal amino acid sequence by use of a spinning cup sequenator, or (2) to homogeneity by SDS-PGGE under non-reducing or reducing conditions using Coomassie blue or, preferably, silver stain. Isolated polypeptide includes

10

15

20

25

30

35

polypeptide *in situ* within recombinant cells, since at least one component of the CDK4 binding peptide natural environment will not be present. Ordinarily, however, isolated polypeptide is prepared by at least one purification step.

An "isolated" nucleic acid molecule is a nucleic acid molecule that is identified and separated from at least one contaminant nucleic acid molecule with which it is ordinarily associated. An isolated nucleic acid molecule is other than in the form or setting in which it is found in nature. However, an isolated nucleic acid molecule includes nucleic acid molecules contains in cells that ordinarily express the protein encoded by that nucleic acid molecule where, for example, the nucleic acid molecule is in a chromosomal location different from that found in nature.

The term "control sequence" refers to DNA sequences necessary for the expression of an operably linked coding sequence in a particular host organism. The control sequences that are suitable for prokaryotes, for example, include a promoter, optionally an operator sequence, and a ribosome binding site. Eukaryotic cells are known to utilize promoters, polyadenylation signals, and enhancers.

Nucleic acid is "operably linked" when it is placed into a functional relationship with another nucleic acid sequence. For example, DNA for a Nuclear Localization Signal is operably linked to DNA encoding the CDK4 binding peptide if the expressed protein is localized in the nucleus thereby; a promoter or enhancer is operably linked to a coding sequence if it affects the transcription of the sequence. Generally, "operably linked" means that the DNA sequences being linked are contiguous, preferably in the same reading frame. However, enhancers do not have to be contiguous. Linking is accomplished by ligation at convenient restriction sites. If such sites do not exist, a synthetic oligonucleotide adaptor or linkers can be used in accordance with conventional practice.

The term "antibody" is used in the broadest sense and specifically covers single anti-CDK4 binding peptide monoclonal antibodies (including agonist, antagonist, and neutralizing or blocking antibodies) and antibody compositions with polyepitopic specificity. The term "monoclonal antibody" as used herein refers to an antibody obtained from a population of substantially homogeneous antibodies, i.e., the individual antibodies comprising the population are identical except for possible naturally occuring mutations that may be present in minor amounts.

"Active" or "activity" for the purposes herein refers to form(s) of the CDK4 binding peptide which retain the biologic activities of native or naturally occurring MyoD having a CDK4 binding domain.

The terms "cancer" and "cancerous" refer to or describe the physiological condition in mammals that is typically characterized by unregulated cell growth. Examples of cancer include but are not limited to, carcinoma, lymphoma, blastoma, sarcoma, and leukemia. Particular examples of cancer include osteosarcoma and rhabdomyosarcoma.

The terms "treating", "treatment", and "therapy" as used herein refer to curative therapy, prophylactic therapy and preventative therapy.

Amino acids may be classified according to the polarities of their side chains. An amino acid having a "non-polar side chain" refers to an amino acid having a side chain that is generally hydrophobic and tends to cluster with other hydrophobic molecules when placed in an aqueous environment, such as water. Examples of amino acids having non-polar side chains are glycine, alanine, valine, leucine, isoleucine, methionine, proline, phenylalanine, and tryptophan. An amino acid having an "uncharged polar side chain" refers to an amino acid having a side chain which includes a hydroxyl, amide or thiol group. Examples of amino acids with uncharged polar side chains are serine, threonine, asparagine, glutamine, tyrosine and cysteine. An amino acid having a "charged polar side chain" refers to an amino acid having a side chain that is generally hydrophyllic. Examples of amino acids having charged polar side chains include lysine, arginine, histidine, aspartic acid and glutamic acid. An amino acid having an "uncharged side chain" includes amino acids with both "non-polar side chains" and "uncharged polar side chains." An amino acid having a "polar side chain" includes amino acids with both "uncharged polar side chains" and "charged polar side chains." Other non-standard amino acids may also be classified using this system.

25

30

35

5

10

15

20

#### Composition

The invention is directed towards a newly identified and isolated amino acid sequences and a nucleic acid sequence coding therefore. The isolated peptide of the invention is capable of specifically binding cyclin dependent kinase (CDK4), thereby inhibiting the activity of CDK4 and cell growth. The isolated peptide will hereinafter be referred to as a "CDK4 binding peptide." The CDK4 binding peptide of the invention includes the following fifteen amino acid sequence: Tyr-Ser-Gly-Pro-Pro-Xaa<sub>1</sub>-Xaa<sub>2</sub>-Xaa<sub>3</sub>-Arg-Arg-Xaa<sub>4</sub>-Asn-Xaa<sub>5</sub>-Tyr-Xaa<sub>6</sub> [SEQ. ID NO: 1] wherein Xaa<sub>1</sub> is an amino acid having an uncharged polar side chain, including Cys or Ser; Xaa<sub>2</sub> is an amino acid having an uncharged side chain, including Ser or

10

15

20

25

30

Gly; Xaa<sub>3</sub> is an amino acid having an uncharged side chain, including Ser, Ala or Pro; Xaa<sub>4</sub> is an amino acid having a polar side chain, including Arg or Gln; Xaa<sub>5</sub> is an amino acid having an uncharged side chain, including Ser, Cys or Gly; and Xaa<sub>6</sub> is an amino acid having a charged polar side chain, including Asp or Glu. Examples of amino acid sequences which correspond to the CDK4 binding peptide of this invention are depicted in Figure 14.

The invention is also directed towards a nucleic acid sequence encoding the CDK4 binding peptide sequence. The nucleotide sequence which encodes the 15 amino acid CDK4 binding peptide of the claimed invention can be determined from various published nucleotide sequences for MyoD. See, Table 1. For example, (a) the complete nucleotide sequence of human MyoD can be found in GenBank at Accession Number X56677 or EMBL Accession Number AF027148 which correspond to Pearson-White, S.H., "Human MyoD: cDNA and deduced amino acid sequence" Nucleic Acids Res. 19(5): 1148 (1991) and Chen, B., et al. "Methylation alterations of the MyoD1 upstream region are predictive of subclassification of human rhabdosarcomas" Am J. Pathol. 152(4): 1071-1079 (1998); (b) the nucleotide sequence which encodes chicken MyoD can be found in GenBank at Accession Number X16189 which corresponds to Lin et al., "An avian muscle factor related to MyoD1 activates muscle-specific promoters in nonmuscle cells of different germ-layer origin and in BrdU-treated myoblasts," Genes Dev., 3(7):986-96 (1989); (c) the nucleotide sequence which encodes Zebrafish MyoD can be found in GenBank/EMBL at Accession Number Z36945 which corresponds to Weinberg, et al. "Development regulation of zebrafish MyoD in wild-type, no tail and spadetail embryos" Development 122:2711-280 (1996). It should be noted that the amino acids which correspond to the CDK4 binding peptide for the various organisms provided above are not the same for each.

Applicants also recognize, and include within the scope of their invention, a nucleic acid sequence for the CDK4 binding peptide which contains codons that are modified according to optimal codon frequencies for a particular cellular host. Redundancy in the genetic code permits variation in the gene sequence. In particular, specific codon preferences are recognized for a specific host such that the disclosed sequence can be adapted as preferred for the desired host. For example, rare codons having a frequency of less than about 20% in known sequences of the desired host are preferably replaced with higher frequency codons. A standard

textbook on cell biology such as Genes V, Benjamin Lewin ed. Oxford University Press (1994) (see, Chapter 7, page 172, Fig. 7.9) can be consulted for information on the degeneracy of the genetic code and to provide information regarding the various nucleic acid codons and to which amino acid they correspond.

## 5 Table 1.

Source/Species	GenBank/EMBL	Journal
	Accession No.	
Chicken	L34006	Dechesne, C.A., et al. "E-box and MEF-2 independent muscle-specific expression, positive autoregulation, and cross-activation of the chicken MyoD (CMD1) promoter reveal an indirect regulatory pathway" Mol. Cell. Biol. 14(8): 5474-86 (1994)
Human	X5667; AF027148 (EMBL)	Pearson—White, S.H., "Human MyoD: cDNA and deduced amino acid sequence" Nucleic Acids Res. 19(5): 1148 (1991); Chen, B., et al. "Methylation alterations of the MyoD1 upstream region are predictive of subclassification of human rhabdosarcomas" Am J. Pathol. 152(4): 1071–1079 (1998).
Rat	M84176 (EMBL)	Vaida, T.B., et al. "Isolation and structural analysis of the Rat MyoD gene" Gene 116: 223–230 (1992)
Murine	M18779; M84918;	Davis, R.L., et al "Expression of a single transfected cDNA converts fibroblasts to myoblasts" Cell 51: 987–1000 (1987); Pinney, D.F., et al. "Myogenic lineage determination and differentiation: Evidence for a regulatory gene pathway" Cell 53: 781–93 (1988)
Pig	U12574	Chang, K.C., et al. "Cloning and in vivo expression of the pig MyoD gene" J. Muscle Res. Cell. Motil. 16(3): 243–47 (1995).
Xenopus-A	X16106/M31116	Hopwood, N.D., et al. "MyoD expression in the forming somites is an early response to mesoderm induction in Xenopus embryos" EMBO J. 8(11): 3409–17 (1989); Scales, J.B., et al. "Two distinct Xenopus genes with homology to MyoD1 are expressed before somite formation in early embryogenesis" Mol. Cell. Biol. 10: 1516–1524 (1990); and
Zebrafish	Z36945	Weinberg, et al. "Developmental regulation of zebrafish MyoD in wild—type, no tail and spadetail embryos" Development 122: 2711–280 (1996).

10

15

20

25

30

The peptide of the invention is an amino acid domain from the carboxyterminal region of muscle specific transcription factor (MyoD) which includes the CDK4 binding domain. The CDK4 binding peptide sequence of the invention is well conserved among MyoD proteins of the higher vertebrates (Figure 14). While not wishing to be bound by theory, the inventors believe that the peptide sequence of the invention specifically binds to CDK4 to inhibit kinase activity. In particular, the inventors believe that interaction inhibits the kinase activity of CDK4 towards retinoblastoma protein (RB). This interaction provides an alternative mechanism that allows MyoD to restrict cell differentiation, independent of a direct interaction with the retinoblastoma protein.

Unlike other vertebrate myogenic factors, the interaction between the peptide of the invention and CDK4 does not involve a highly conserved bHLH domain. Helix-Loop-Helix (HLH) is a motif that is found in a large family of nuclear proteins which appear to play a fundamental role in cellular growth, transcription, and differentiation. The structure, function and location of the bHLH domain is known to those of skill in the art and is published by Ma et al, Cell, 77: 451–459; (1994). Additionally, Figure 9 provides the amino acid locations of the various domains of chicken MyoD.

In contrast to previous studies that only correlate the bHLH domain of MyoD with growth arrest (Crescenzi et al., <u>PNAS USA</u>, 1990, 87(21):8442–6; and Sorrentino et al., <u>Nature</u>, 1990 345(6278):813–5), the inventors have found that the amino terminus of MyoD exhibits no growth arrest properties, whereas the bHLH domain, and to a greater extent, the carboxy terminus of MyoD both inhibit cell growth. In fact, the inventors have found that the CDK4 binding peptide alone is sufficient to inhibit cell growth and is actually the most efficient growth inhibitory domain in MyoD.

#### **Nuclear Localization Signal**

The inventors have also discovered that cyclin Dl regulates MyoD function and the onset of myogenesis by controlling the cellular location of CDK4. In the presence of cyclin D1, CDK4 is translocated from the cytoplasm to nucleus. If it is in sufficient excess to MyoD, nuclear CDK4 sequesters nuclear MyoD and thus inhibits activation of muscle specific genes.

Correspondingly, the inventors have found that the growth inhibitory effect of the CDK4 binding peptide of the invention is enhanced when the peptide further includes nuclear localization signal (NLS) (data not shown). The NLS can be placed anywhere in the protein. For example, the NLS signal can be attached at the amino terminus of the fifteen amino acid MyoD CDK4 binding peptide and fused to GFP. Suitable NLS are described by Fischer–Fantuzzi, et al., Mol. Cell. Biol.; 8: 5494–5503 (1988).

## **Amino Acid Variants**

Covalent modifications of the CDK4 binding peptide are included within the scope of this invention. One type of modification includes deamidation of glutaminyl and asparaginyl residues to the corresponding glutamyl and arpartyl residues, respectively, hydroxylation of proline and lysine, phosphorylation of hydroxyl groups of seryl or threonyl residues, methylation of the  $\alpha$ -amino groups of lysine, arginine and histidine side chains, acetylation of the N-terminal amine, and amidation of any C-terminal carboxyl groups. Another covalent modification includes altering the native glycosylation pattern of the polypeptide, for example, by deleting one or more carbohydrate moieties found in the native CDK4 binding peptide and/or adding one or more glycosylation sites that are not present in the native sequence. Another type of covalent modification includes linking the CDK4 binding peptide to one of a variety of non-proteinaceous polymers, e.g., polyethylene glycol, polypropylene glycol, orpolyoxyalkylenes.

The CDK4 binding peptide of the invention may also be modified to form a chimeric molecule which includes the CDK4 binding peptide fused to another, heterologous peptide or amino acid sequence. The inventors have found that such fusion proteins can induce growth arrest as efficiently as full—sized MyoD. In one embodiment, the chimeric molecule includes the CDK4 binding peptide fused to a nuclear localization signal. Other examples of heterologous peptide or amino acid sequences include peptides which target the molecule to specific cells, for example, by attachment to an antibody against a surface protein of the target cell. Additionally, the protein can be targeted to various compartments of the cell including the nucleus, the mitochondria (Rizutto et al, Nature; 358: 325–327 (1992)), and endoplasmic reticulum (Munro et al, Cell; 48: 899–907 (1987)).

## **Nucleic Acid Variants**

Applicants recognize, and include within the scope of their invention, a nucleotide sequence (DNA or RNA) encoding the peptide sequence which contains codons that are modified according to optimal codon frequencies for a particular cellular host.

Redundancy in the genetic code permits variation in the gene sequence encoding the peptide sequence of the invention. In particular, specific codon preferences are recognized for a specific host such that the disclosed sequence can be adapted as preferred for the desired host. For example, rare codons having a frequency of less than about 20% in known sequences of the desired host are preferably replaced with higher frequency codons.

## Preparation of CDK4 Binding Peptide

The description below relates primarily to production of CDK4 binding peptide by culturing cells transformed or transfected with a vector containing CDK4 nucleic acid. It is, of course, contemplated that alternative methods, which are well known in the art, may be employed to prepare CDK4 binding peptide. For instance, the peptide, or portions thereof, may be produced by direct peptide synthesis using solid—phase techniques. *In vitro* protein synthesis may be performed using manual techniques or by automation. Automated synthesis may be accomplished, for example, using an Applied Biosystems Peptide Synthesizer (Foster City, CA) following the manufacturer's instructions. Various portions of the CDK4 binding peptide may be chemically synthesized separately and combined using chemical or enzymatic methods.

25

30

5

10

15

20

#### Isolation of DNA Encoding the CDK4 Binding Peptide

DNA encoding the CDK4 Binding Peptide may be obtained from a cDNA library. The library can be screened with probes (such as antibodies to the CDK4 binding peptide or oligonucleotides) designed to identify the gene of interest or the protein encoding by it. Screening the cDNA or genomic library with the selected probe may be conducted using procedures known to those of skill in the art.

## Selection and Transformation of Host Cells

Host cells may be transfected or transformed with expression or cloning vectors described herein for production of the CDK4 binding peptide. The host cells are cultured in conventional nutrient media modified as appropriate for inducing promoters, selecting transformants, or amplifying the gene encoding the desired sequences. The culture conditions, such as media, temperature, pH and the like, can be selected by skilled artisans without undue experimentation.

Methods of transfection are known to ordinarily skilled artisans, for example, CaPO<sub>4</sub> and electroporation. Alternately, the nucleic acid sequence can be introduced into viable cells using liposomes, microinjection, cell fusion, DEAE-dextran. Depending on the host cell used, transformation is performed using standard techniques appropriate to such cells.

Suitable host cells for cloning or expressing the DNA in the vectors herein include

prokaryote, yeast, or higher eukaryotic cells. Suitable prokaryotes include, but are not limited to eubacteria, such as Gram-negative or Gram-positive organisms, for example, Enterobacteriaceae such as E. Coli. Eukaryotic cells, including, but not limited to Sf9 cells, 10T1/2 fibroblasts and C2C12 myoblast can also be transfected.

## 20 <u>Vectors</u>

5

10

15

25

30

35

The nucleic acid sequence encoding the CDK4 binding peptide may be inserted into a replicable vector for cloning (amplification of the DNA) or for expression. Various vectors are publicly available. The vector may, for example, be in the form of a plasmid, cosmid, viral particle, phage, or, a baculovirus vector. The appropriate nucleic acid sequence may be inserted into the vector by a variety of procedures. In general, DNA is inserted into an appropriate restriction endonuclease site(s) using techniques known in the art. Vector components generally include, but are not limited to, one or more of a signal sequence, an origin of replication, one or more marker genes, an enhancer element, a promoter, and a transcription termination sequence. Construction of suitable vectors containing one or more of these components employs standard ligation techniques which are known to skilled artisans.

The CDK4 binding peptide may be produced recombinantly not only directly, but also as a fusion polypeptide which a heterologous polypeptide, for example, a nuclear localization signal.

Expression and cloning vectors typically include a selection gene, also termed a selectable marker. Typical selection genes encoding proteins that (a) confer resistance to antibiotics or other toxins, e.g., ampicillin, neomycin, methotrexate, or tetracycline, (b) complement auxotrophic deficiencies, or (c) supply critical nutrients not available from complex media.

Expression and cloning vectors also typically include a promoter operably linked to the nucleic acid sequence which encodes the CDK4 binding peptide. Promoters recognized by a variety of potential host cells are well known, and include inducible and non–inducible promoters.

10

15

20

5

#### **Detecting Gene Amplification/Expression**

Gene amplification and/or expression may be measured in a sample directly, for example, by conventional Southern blotting, Northern blotting to quantify the transcription of mRNA, dot blotting (DNA analysis), or *in situ* hybridization using an appropriately labeled probe based on the sequence provided herein. Alternately, gene expression may be measured by immunological methods, such as immunohistochemical staining of cells or tissue sections and assay of cell culture or body fluids, to quantify directly the expression of a gene product. Antibodies useful for immunohistochemical staining and/or assay of sample fluids may be either monoclonal or polyclonal, and may be prepared in any mammal. Conveniently, the antibodies may be prepared against a native sequence of the CDK4 binding peptide or against a synthetic peptide based on the sequences provided herein or against an exogenous sequence fused to the CDK4 binding peptide and encoding a specific antibody epitope.

25

30

#### **Purification**

CDK4 binding peptide may be recovered from culture medium or from host cell lysates, for example, by fractionation on an ion-exchange column; ethanol precipitation; reverse phase HPLC; chromatography on silica or on a cation-exchange resin such as DEAE; chromatofocusing; SDS-PAGE; ammonium sulfate precipitation; gel filtration using, for example, Sephadex G-75; protein A Sepharose columns; and metal chelating columns. Such methods of protein purification are known to those of skill in the art. The purification step selected will depend, for example, on the nature of the production process used.

10

15

20

25

30

Nucleotide sequences (or their complement) encoding the CDK4 binding peptide have various applications in the art of molecular biology, including uses as hybridization probes, in chromosome and gene mapping and in the generation of anti—sense RNA and DNA. Nucleotide sequences encoding the CDK4 binding peptide are also useful for the preparation of CDK4 binding peptide by the recombinant techniques described herein. The nucleic acid sequences are also useful to generate transgenic animals or "knock out" animals which, in turn, are useful in the development and screening of therapeutically useful reagents. A transgenic animal is an animal having cells that contain a transgene, which transgene was introduced into the animal or an ancestor of the animal at a prenatal, or embryonic, stage. A transgene is a DNA which is integrated into the genome of a cell from which a transgenic animal develops.

The CDK4 binding peptide, as disclosed in the present invention, can be employed therapeutically to inhibit cell proliferation, CDK4 kinase activity or to treat solid tumors, particularly those which over—express CDK4.

The CDK4 binding peptide can be used in *in vivo* or *ex vivo* gene therapy methods. In gene therapy applications, genes are introduced into cells in order to achieve *in vivo* synthesis of a therapeutically effective gene product, for example, the replacement of a defective gene. "Gene therapy" includes both conventional gene therapy where a lasting effect is achieved by a single treatment, and in the administration of gene therapeutic agents, which involves the one time or repeated administration of a therapeutically effective DNA or mRNA. Antisense RNAs and DNAs can be used as therapeutic agents for blocking the expression certain genes *in vivo*.

Methods of transfection are known to ordinarily skilled artisans. The techniques vary depending upon whether the nucleic acid is transferred into cultured cells *in vitro* or *in vivo*, in the cells of the intended host. Examples of suitable techniques for introducing the nucleic acid sequence can be into viable include the use of liposomes, microinjection, cell fusion, DEAE—dextran, calcium phosphate precipitation and electroporation. The currently preferred *in vivo* gene transfer techniques include transfection with viral (typically retroviral) vectors Depending on the host cell used, transformation is performed using standard techniques appropriate to such cells.

The CDK4 binding peptide and variants thereof may be used therapeutically to inhibit the activity of CDK4 and cell growth or cell division. According to the invention, a composition which includes the peptide of the invention is administered to a cell culture or a patient in need of such treatment wherein the composition includes an effective amount of the CDK4 binding peptide of the invention. The inhibition of CDK4 activity and/or cell growth has applications in diseases in which CDK4 is over expressed, for example, cancer. In particular, the CDK4 binding peptide can be administered to treat solid tumors which over—express CDK4, such as rhabdomyosarcoma (RMS) and osteosarcoma.

10

15

20

25

30

5

## Pharmaceutical Composition

Another aspect of the invention is directed towards a pharmaceutical composition for inhibiting CDK4 kinase activity, inhibiting cell division, or treating solid tumors which over—express CDK4. The pharmaceutical composition of the invention includes an effective amount of the CDK4 binding peptide of the invention and a pharmaceutically acceptable excipient or carrier. Suitable pharmaceutical carriers are known by those of skill in the art. Typically, an appropriate amount of a pharmaceutically acceptable salt is used in the formulation to render the formulation isotonic. Examples of the carrier include buffers such as saline, Ringer's solution and dextrose solution. The pH of the solution is preferably from about 5 to about 8, and more preferably from about 7.4 to about 7.8. Further carriers include sustained release preparations such as semipermeable matrices of solid hydrophobic polymers, which matrices are in the form of shaped articles, e.g., films, liposomes, or microparticles. It will be apparent to those persons skilled in the art that certain carriers may be more preferable depending upon, for instance, the route of administration and concentration of the CDK4 binding peptide being administered.

#### **Delivery**

Various methods of administration of peptides are known to those skilled in the art. Such methods of administration may include, but are not limited to, surface application, oral and parenteral routes, subcutaneous injection, intravenous injection or other pharmaceutical methods of delivery. The peptide can be targeted to specific cells for use in the treatment of cancer. For example, the peptide can be delivered by liposomes, lipophilic tag, injection, viral vectors or fused to a cell specific antibody.

35 Variants too.

15

20

25

30

Appropriate dosages of the peptides of the invention will depend upon the condition presented by the individual subject. The dosage administered will vary depending upon factors such as the pharmacodynamic characteristics of the particular agent, and its mode and route of administration; age, health and weight of the recipient; nature and extent of symptoms, kind of concurrent treatment, frequency of treatment, and the effect desired. Effective dosages and schedules for administration may be determined empirically, and making such determinations is within the skill in the art.

## 10 Anti-CDK4 Binding Peptide Antibodies

The invention further provides anti-CDK4 binding peptide antibodies. Exemplary antibodies include polyclonal, monoclonal, humanized, bispecific and heteroconjugate antibodies.

The anti-CDK4 binding peptide antibodies may include a polyclonal antibody. Method of preparing polyclonal antibodies are known to those skilled in the art. Polyclonal antibodies can be raised in a mammal, for example, by one or more injections of an immunizing agent and, if desired, an adjuvant. Typically, the immunizing agent and/or adjuvant will be injected in the mammal by multiple subcutaneous or intraperitoneal injections. The immunizing agent may include the CDK4 binding peptide, and variants or fusion proteins thereof.

Alternatively, the anti-CDK4 binding peptide antibodies may be monoclonal antibodies. Monoclonal antibodies may be prepared using hybridoma methods which are known to those of skill in the art. In a hybridoma method, a mouse, hamster, or other appropriate host animal is typically immunized with an immunizing agent to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the immunizing agent. The immunizing agent may include the CDK4 binding peptide, and variants or fusion proteins thereof. Alternately, the lymphocytes may be immunized *in vitro*. The lymphocytes are then fused with an immortalized cell line using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell. The culture medium in which the hybridoma cells are cultured can then be assayed for the presence of monoclonal antibodies directed against the CDK4 binding peptide, for example, by immunoprecipitation or by an *in vitro* binding assay, such as radioimmunoassay (RIA) or enzyme linked immunoabsorbent assay (ELISA). Such techniques are

35 known in the art.

10

15

20

25

30

35

The antibodies may be monovalent antibodies. Methods for preparing monovalent antibodies are well known in the art. For example, one method involves recombinant expression of immunoglobulin light chain and modified heavy chain. The heavy chain is truncated generally at any point in the Fc region so as to prevent heavy chain crosslinking. Alternatively, the relevant cysteine residues are substituted with another amino acid residue or are deleted to prevent crosslinking. *In vitro* methods are also suitable for preparing monovalent antibodies. Digestion of antibodies to produce fragments thereof, particularly, Fab fragments, can be accomplished using routine techniques known in the art.

The anti-CDK4 binding peptide antibodies of the invention may further comprise humanized antibodies or human antibodies. Humanized forms of nonhuman (e.g., murine) antibodies are chimeric immunoglobulins, immunoglobulin chains or fragments thereof (such as Fv, Fab, Fab', F(ab')<sub>2</sub> or other antigen-binding subsequences of antibodies) which contain minimal sequence derived from nonhuman immunoglobulin. Humanized antibodies include human immunoglobulins (recipient antibody) in which residues from a complementary determining region (CDR) of the recipient are replaced by residues from a CDR of a non-human species (donor antibody) such as a mouse, rat or rabbit having the desired specificity, affinity or capacity. In some instances, Fv framework residues of the human immunoglobulin are replaced by corresponding non-human residues. Humanized antibodies may also comprise residues which are found neither in the recipient antibody nor in the imported CDR or framework sequences. In general, the humanized antibody will comprise substantially all of at least on, and typically two, variable domains, win which all or substantially all of the CDR regions correspond to those of a non-human immunoglobulin and all or substantially all of the FR regions are those of a human immunoglobulin consensus sequence. The humanized antibody optimally also comprises at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin. Methods for humanizing non-human antibodies are well known in the art.

Bispecific antibodies are monoclonal, preferably human or humanized, antibodies that have binding specificities for at least two different antigens. Heteroconjugate antibodies are composed of two covalently joined antibodies.

The anti-CDK4 binding peptide antibodies of the invention have various utilities. The antibodies can be used in diagnostic assays for CDK4 binding peptide, e.g., detecting its expression in specific cells, tissues serum or tumors. Various

10

15

20

30

35

diagnostic assay techniques are known in the art and may be used, such as competitive binding assays, direct or indirect sandwich assays and immunoprecipitation assays conducted in either heterogeneous or homogeneous phases. The antibodies used in the diagnostic assays can be labeled with a detectable moiety capable of producing, either directly or indirectly, a detectable signal. For example, the detectable moiety may be a radioisotope (e.g., <sup>3</sup>H, <sup>14</sup>C, <sup>32</sup>P, <sup>35</sup>S, or <sup>125</sup>I), a fluorescent or chemiluminescent compound (e.g., fluorescein isothiocyanate, rhodamine, or luciferin) or an enzyme (e.g., alkaline phosphatase, beta–galactosidase, or horseradish peroxidase). Any known method for conjugating the antibody to the detectable moiety may be employed.

The antibodies of the invention are also useful for affinity purification of CDK4 binding peptide from recombinant cell culture or natural sources. In this process, the antibodies are immobilized on a suitable support, such as Sephadex resin or filter paper, using methods well known in the art. The immobilized antibody is then contacted with a sample containing CDK4 binding peptide to be purified, and thereafter the support is washed with a suitable solvent that will remove substantially all the material in the sample except the CDK4 binding peptide, which is bound to the immobilized antibody. Finally, the support is washed with another suitable solvent that will release the CDK4 from the antibody. The antibodies also have therapeutic utility.

All references cited in the specification, including in the Working Examples, are hereby incorporated by reference herein.

#### **WORKING EXAMPLES**

The following examples are provided to more fully illustrate the principles and practices of the invention. The examples are not intended in any way to limit the scope of the invention.

## I. In vitro Examination of MyoD-CDK4 Interaction

Cloning, Cell Culture and Preparation of Cell Extract

Using primers with the corresponding restriction sites, chicken MyoD, myogenin, and E12 were cloned into EcoRl/Hind3 digested and CIP-treated pM and pVP16 vectors (Clonetech) using standard PCR methods. Sf9 cells were coinfected with baculoviral stocks encoding CDK4 and cyclin to provide a source of activated CDK4 by methods known to those of skill in the art. After incubation for 48 hours

at 23° C, cell extracts were prepared from the coinfected Sf9 cells using standard methods in the art.

## **Binding Assay**

5

10

15

20

25

The Sf9 cell extract was incubated with agarasoe bound with either glutathione transferase (GST) alone, or GST fusion proteins encoding one of the four different vertebrate myogenic factors from chicken: CMD1 (MyoD), Ch-myogenin, Ch-myf5 an Ch-mrf4 using methods known to those of skill in the art. Any GST complexes formed were recovered on glutathione-Sepharose beads. The recovered GST complexes were then washed, heated in sample buffer and loaded on an SDS-PAGE gel.

### Western Blot

The gel was then analyzed by immunoblot with anti-CDK4 affinity purified rabbit polyclonal antibody on HyBond ECL (Amersham). The anti-CKD4 antibody was diluted 1:2,000-5,000. The second antibody was HRP anti-rabbit and was diluted 1:10,000 (Pierce cat. no. 31460). The signal was detected using SuperSignal Ultra Reagent (Pierce cat. no. 34075).

The results indicate that CDK4 binds specifically to GST-MyoD. Only the GST-MyoD bound CDK4 (data not shown). Binding assays performed with extracts from infected Sf9 cells containing only CDK4 or with bacterially produced CDK4 gave the same results (data not shown). Additionally, the ubiquitously expressed bHLH factors E12 and E47 (proteins that form active heterodimers with MyoD) did not bind CDK4 as GST fusion proteins (data not shown). This demonstrates that active kinase is not required for CDK4 to bind specifically to MyoD in vitro. Furthermore, the interaction is not mediated by an unknown protein or proteins in the Sf9 cell extracts since purified bacterial proteins give identical results.

## 30 II. In vivo Determination of MyoD-CDK4 Interaction

Three approaches were used to determine whether or not the observed MyoD-CDK4 interaction occurred <u>in vivo</u>. The combined results from these approaches demonstrate that MyoD and CDK4 interact specifically <u>in vivo</u>.

## A. Coimmunoprecipitation

#### **Transfection**

Vectors were prepared essentially as described in Example I. Sf9 cells were coinfected to produce either MyoD and CDK4 or MyoD and cyclin Dl protein combinations. After incubating the coinfected Sf9 cells for 48 hours at 23° C, cell extracts were prepared essentially as described in Example 1.

## 10 <u>Coimmunoprecipitation</u>

Coimmunoprecipitation reactions were performed sequentially using the Sf9 cell extracts (adjusted for equal protein concentrations in each reaction), under sequential low (no SDS) and high (with SDS) stringency conditions with the antibody combinations shown in Figure 1 using antibody dilutions of 1:200.

Immunoblotting

Secondary immune complexes were resuspended directly in SDS loading buffer and after SDS-PAGE, proteins were detected by immunoblot essentially as described in Example I, above.

Results

The results are shown in Figure 1: cells coinfected with baculovirus encoding MyoD and cyclin Dl are in lanes 1 to 3; MyoD and CDK4 are in lanes 4 and 5; and MyoD, CDK4, and cyclin Dl are in lanes 6 to 11. A comparison of lanes 3 and 5 shows that MyoD interacted with CDK4 but not cyclin Dl. A comparison of lanes 9 and 11 shows that coexpression of MyoD, CDK4 and cyclin Dl proteins results in MyoD formation of immune complexes containing both CDK4 and cyclin Dl. This shows that MyoD binding to the cyclin Dl—CDK4 complex does not dissociate the cyclin from the kinase.

MyoD expressed in Sf9 cells is phosphorylated on serine residues, as is MyoD expressed in primary cultures of embryonic chick breast muscle. Because MyoD does not readily form homodimers <u>in vitro</u> unless it is dephosphorylated, MyoD is assumed to bind to CDK4 as a monomer in the immunoprecipitation reactions from Sf9 cell extracts.

20

25

30

15

25

Coimmunoprecipitation reactions substituting either virally expressed myf5 or myogenin for MyoD failed to produce immune complexes with CDK4, in agreement with <u>in vitro</u> GST fusion-binding results (data not shown).

## 5 B. <u>Mammalian Two-Hybrid System</u>

The interaction between MyoD and CDK4 in vivo was examined using a mammalian version of the yeast two hybrid system described by Finkel et al., (1993) <u>J Biol Chem</u> 268, 5–8.

## 10 <u>Vector Preparation</u>

MyoD and myogenin (MGN) were each cloned into the gal4 DNA-binding domain vector, pM. CDK4 was inserted into the vpl6 activation domain plasmid, pVP16 using methods known to those of skill in the art.

## 15 <u>Transfection</u>

10T1/2 fibroblasts (3xl0<sup>5</sup> cells per 60mm dish or multiwell plate) grown in DMEM with either 10% fetal calf serum were co-transfected with either gal4–MyoD or gal4–MGN along with vpl6–CDK4 and a luciferase reporter using Fugene–6 (Boehringer–Mannheim) according to the manufacturer's directions. 1 μg gal4 luciferase reporter, 2 μg each of the gal4 (pM vector, Clonetech) and vpl6 (pvpl6 vector, Clonetech) constructs, and 300 ng of the beta galactosidase reporter pCH110 (Pharmacia), a total of 5.3 μg DNA, were used in each transfection. The pSV2–beta galactosidase reporter was used to normalize transfection efficiencies.

After 24–36 hours, if required, cells were placed in 2% horse serum plus insulin (10  $\mu$ g/ml) to induce differentiation and destabilize cyclin Dl. 24–48 hrs later cells were harvested for a luciferase reporter assay.

#### Luciferase Assay

The gal4 luciferase reporter was constructed by exchanging the CAT cassette from pG5 (Clonetech) for the luciferase cassette in pGL3 (Promega). To do so, pG5 was cut with Eco R1 and flush ended with T4 polymerase. The luciferase gene was digested with BamH1/Hind3, flush ended with T4 polymerase and inserted into pG5. The transfection was performed using Fugene-6 (Boehringer-Mannheim) according to the manufacturer's directions.

24-48 hours after transfection, the luciferase reporter assay was performed according to the manufacturer's directions (Promega, cat. no. E4030) and measured in a Victor 1420 Multilabel Counter. The assay was repeated at least three times with similar results. The luciferase reporter assay was normalized for transfection efficiency with a cotransfected beta galactosidase reporter and by total protein in the cell extract with similar results.

### Results

5

10

15

20

25

MyoD interactions with CDK4 are shown in lanes 1 to 5 and myogenin interactions with CDK4 are shown in lanes 6-10. Gal4 MyoD-vp16 E12 and Gal4 MGN-vp16 E12 interactions (previously established heterodimer partner combinations) were used a positive controls. See, Figure 2, lanes 1 to 5. Despite the high cellular levels of endogenous CDK4 competing in the two hybrid reaction, a four-fold increase in vpl6 CDK4-dependent luciferase reporter activity was routinely detected (an average of three experiments) with the use of the gal4-MyoD construct (Figure 2, lanes 2 and 3) and not with gal4-MGN (Figure 2, lanes 7 and 8).

## C. Immunoprecipitation

An immunoprecipitation assay was performed to determine whether MyoD and CDK4 interact in dividing C2C12 myoblasts.

# Labeling and Preparation of Nuclear Extracts

C2C12 myoblasts were maintained in growth medium and labeled with <sup>35</sup>S (90 minute labeling) using methods known to those of skill in the art. Nuclear extracts were prepared from <sup>35</sup>S-labeled cultures (90 minute labeling) of C2C12 myoblasts maintained in growth medium.

#### Coimmunoprecipitation

Nuclear extracts containing equal counts were sequentially
coimmunoprecipitated under sequential low (no SDS) and high (with SDS)
stringency conditions with indicated primary and secondary antibody combinations
shown in Figure 3 using essentially the same protocol as that described in Example
II.

## Results

5

15

20

25

35

MyoD-CDK4 immune complexes were identified in dividing myoblasts (Figure 3, lane 6). However, such complexes were not detected in differentiated muscle cultures (data not shown). MyoD and CDK4 interact in dividing C2C12 myoblasts. Lane 6 (Figure 3) shows that anti-MyoD pulls down CDK4 in dividing myoblasts that can be detected with the secondary CDK4 antibody.

## III. Band Shift Assays

To ascertain whether or not CDK4 interferes directly with MyoD transactivation functions, the DNA-binding activity of both MyoD homodimers as well as MyoD-E12 heterodimers was measured in band-shift assays using the bacterially expressed and histidine-tag purified proteins, as described by Shirakata et al., *Genes Dev* 7, 2456-70 (1993).

## Cloning and Transfection

His-tagged CDK4 was cloned as the BamHl/Hind3 fragment in the pRSET plasmid (Invitrogen). Chicken MyoD, myogenin, and E12 were cloned into EcoRl/Hind3 digested and CIP-treated pM and pVP16 vectors (Clonetech) using standard PCR methods with primers having the corresponding restriction sites. All constructed clones were checked by sequence analysis on a Perkin-Elmer 310 Genetic Analyzer. His-tagged CDK4 was expressed in the E. Coli strain BL21 as described by Shirakata et al., (1993) Genes Dev., 7:2456-70.

## **Histidine Tag Purification**

The protocol for the histidine tag purification technique is identical to Shirakata, <u>supra</u>, with slight changes that are not critical to the outcome.

#### Renaturation

The his-tagged CDK4 was renatured through sequential dialysis from urea as described by Kato et al., (1994) Mol. Cell. Biol. 14:2713-21.

## **Band Shift Assay**

Band shift assays were performed using the bacterially expressed and histidine-tag purified proteins. The MyoD (10 ng), myogenin (10 ng) and E12 (30 ng) proteins from chicken were used with increasing two-fold increments (50-200

ng) of mouse CDK4 protein in the band shift reaction. Limiting amounts of E12 protein were used in the MyoD band shift reactions to observe any CDK4 effects on both the MyoD homodimer as well as the MyoD-E12 heterodimer. DNA binding by a myogenin-E12 heterodimer was used as a control since myogenin does not bind CDK4 in vitro.

#### Results

A CDK4 concentration dependent inhibition of DNA binding was observed with the MyoD homo— and heterodimer complexes (Figure 4, lanes 1 to 4). Thus, CDK4 protein inhibited DNA—binding by both the MyoD—E12 heterodimer and the MyoD. No CDK4 inhibition of DNA—binding was observed with the myogenin—E12 complex using the same final level (50–200 ng) of CDK4 protein in the shift reaction (Figure 4, lanes 5 to 8). Thus, DNA—binding by the myogenin—E12 heterodimer and the myogenin homodimer were unaffected at the equivalent concentrations of CDK4 protein in the binding reaction.

Because the actual amount of properly renatured CDK4 protein is unknown, the effective inhibitory ratio of CDK4 to MyoD–E12 complex may be significantly less than five–fold. We assume that CDK4 dissociates the DNA–binding complexes, since no evidence for a supershift was noted either with the addition of CDK4 protein or CDK4 antibody to the shift reactions (data not shown). Most importantly, in agreement with the earlier protein binding studies, CDK4 inhibition of DNA–binding by the MyoD–E12 heterodimer did not depend upon cyclin DI and, therefore, does not require the active kinase. Thus, CDK4 specifically inhibits MyoD–E12 DNA binding in the absence of cyclin D1.

25

20

5

10

15

#### IV. Immunofluorescence

Immunofluorescence assays were used to determine whether CDK4 was differentially located in nuclei during myogenesis and to determine the effect of interactions between MyoD, cyclin D1 and CDK4.

30

35

#### <u>Immunofluorescence</u>

To determine whether CDK4 was differentially localized in nuclei during myogenesis, dividing myoblasts in growth medium (GM) as well as differentiated cultures of C2Cl2 mouse myoblasts in differentiation medium (DM) were reacted with antibodies to CDK4 and cyclin D1 and examined by immunofluorescence.

The antibodies to CDK4 and cyclinD1 were used at 1:200-500 dilution (rabbit or mouse anti-CDK4 and anticyclin Dl, Santa Cruz). Second antibody dilutions were also 1:200-500 with either Alexa 488 (green) goat anti-rabbit or mouse, or Alexa 594 (red) goat anti-rabbit or mouse (Molecular Probes).

5

10

15

20

## Results

The results demonstrated that CDK4 is located in the nuclei of dividing myoblasts but translocates to the cytoplasm in differentiated myotubes (data not shown). Furthermore, cyclin Dl is expressed in the nuclei of dividing cells and is completely absent in the nuclei of well differentiated myotubes (data not shown).

## V. <u>Ectopic Expression of Cyclin D1</u>

To determine whether or not ectopic expression of cyclin Dl could relocalize and maintain CDK4 in myotube nuclei and, thus, in dividing myoblast nuclei, an ecdysone inducible cyclin Dl construct was transiently transfected into C2Cl2 myoblasts along with an ecdysone inducible beta galactosidase reporter plasmid.

Because cyclin Dl is rapidly degraded in the absence of growth factors (the conditions used to differentiate muscle cells), a cyclin Dl protein containing an amino acid substitution that renders it resistant to degradation, mutant T286A, was used in place of the wild—type cyclin Dl. Mutant T286A contains an alanine for threonine—286 substitution and is described by Diehl et al., (1997) Genes Dev., 11:957—72.

#### Cloning

25 Cyclin DI was amplified from baculovirus stocks with the appropriate primers and was cloned into the PIND vector (Invitrogen) digested with BamHl/EcoRl.

## **Transfection**

30 C2Cl2 cells were transfected with 2 μg each of pIND/cyclin–Dl(T286A) and pIND/lacZ (Invitrogen). After 24 hours, the cells were switched to differentiation medium (2% horse serum in DMEM with 10 μg/ml insulin).

#### Induction

Cyclin D1(T286A) was induced by the addition of 1 µM of the ecdysone derivative muristerone. 48 hours later, after well-formed myotubes appeared in the cultures, muristerone was added to a final concentration of 1 µM (Invitrogen).

5

10

#### **Immunostaining**

Cells were fixed 24 hours post induction and immunostained for CDK4, cyclin Dl, and lacZ. The first antibodies were used at 1:200–500 dilution (rabbit or mouse anti-CDK4 and anticyclin Dl, Santa Cruz; mouse monoclonal anti-lacZ, Gibco BRL). Second antibody dilutions were also 1:200–500 with either Alexa 488 (green) goat anti-rabbit or mouse, or Alexa 594 (red) goat anti-rabbit or mouse (Molecular Probes) for the immunofluorescent reactions.

#### Results

15

20

25

The results shown induction of both beta galactosidase and stable cyclin Dl (T286A) in well—formed cotransfected myotubes results in the cytoplasmic to nuclear translocalization of endogenous CDK4 (data not shown). There is essentially no detectable cyclin D1 expression in either myotubes or single cells kept in mitogen depleted differentiation medium (data not shown). In contrast, nuclei of rapidly dividing myoblasts maintained in mitogen rich growth medium have high levels of nuclear cyclin D1 (data not shown).

This result clearly demonstrates that the nuclear localization of CDK4 in C2Cl2 myoblasts can be regulated by the cellular level of cyclin Dl, which, in turn, is dependent upon the levels of mitogens in the growth medium. Neither the addition of growth medium to control myotubes cultures nor the muristerone induction of a wild—type cyclin Dl construct resulted in the nuclear translocation of CDK4 (data not shown). Thus, the kinase triggering the ubiquitination and degradation of wildtype cyclin Dl is likely active in myotubes, regardless of the growth conditions.

30

#### VI. Ectopic Expression of CDK4

In this example, the affect of ectopically expressed CDK4 on the transactivation functions of MyoD is Examined.

#### **Plasmid Constructs**

Mouse CDK4 was amplified by PCR from baculovirus stocks as outlined in the MaxBac instruction manual (Invitrogen), with primers containing the corresponding restriction sites and was cloned into pVP16 digested with BamHl/Hind3.

#### **Transfection Protocol**

10T1/2 fibroblasts (3x10<sup>5</sup> cells per 60mm dish or multiwell plate) grown in DMEM with 10% fetal calf serum (growth medium —conditions which promote the nuclear localization of CDK4) were transfected with cotransfected either with an EMSV MyoD or EMSV myogenin expression plasmid, the MCK CAT reporter plasmid, and either an EMSV CDK4 expression construct or the corresponding empty vector using Fugene–6 (Boehringer–Mannheim) according to the manufacturer's directions.

15

20

25

30

10

5

#### Results

Ectopic expression of CDK4 selectively inhibits MyoD transactivation functions in growth medium and the MyoD myogenic conversion of 10T1/2 cells in differentiation medium in the absence of cyclin Dl. CDK4 differentially inhibits MyoD (Figure 5, lanes 3 and 4) and myogenin (Figure 5, lanes 7 and 8) activation of the MCK CAT reporter plasmid in 10T1/2 cells grown in differentiation medium (DM). The ectopic expression of CDK4 was able to significantly suppress the MyoD activation of the MCK–CAT reporter (Figure 5, lanes 3 and 4). In contrast, little suppression was observed on the myogenin–dependent activation of MCK–CAT (Figure 5, lanes 7 and 8). Thus, CDK4 significantly disrupts the activation functions of MyoD, while having a minimal effect on myogenin reporter activation.

Inhibition of myogenic differentiation by cyclin Dl thus appears to be a consequence of the cyclin Dl-dependent nuclear translocation of CDK4 during myogenesis. Furthermore, CDK4 targeted to the myoblast nucleus in the absence of cyclin Dl blocks myogenesis.

#### VII. Nuclear Localization of CDK4

This Example examines whether or not CDK4, in the absence of cyclin Dl, can inhibit myogenesis.

## CDK4 Expression Construct and Transfection

A CDK4 expression construct with a modified with a five-prime nuclear localization signal (NLS) and a three-prime hemagglutinin tag (HA) was prepared. Cultures of 10T1/2 mouse fibroblasts cells were cotransfected as described in Example VI with a fixed amount of pcDNA-MyoD plasmid (1 μg) and increasing amounts of a pcDNA NLSCDK4-HA plasmid (1 to 4μg). Empty pcDNA vector was used as a control and to balance the total plasmid DNA concentration in the transfection. The percentage of myosin positive cells per MyoD positive cell was scored for a minimum of 300 MyoD positive cells for each plasmid concentration. Cultures were maintained in low mitogen differentiation medium to destabilize and eliminate endogenous cyclin Dl and to induce differentiation of the converted cells.

#### Results

Forced nuclear expression of CDK4 inhibits myogenic conversion of 10T1/2 cells in differentiation medium (DM) in the absence of cyclin Dl. Myosin expression in MyoD positive cells was determined as a function of CDK4 plasmid input in the cotransfection. The ratio of myosin positive cells to MyoD positive cells as a function of CDK4 expression plasmid concentration in the transfection was measured. Expression of MyoD and NLSCDK4–HA in the same cell inhibits myogenic conversion (data not shown). MyoD—converted myogenic cells in the absence of NLSCDK4–HA fuse, express MyoD and myosin (data not shown). The forced nuclear localization of CDK4 resulted in an increased number of MyoD—positive single cells that were also negative for myosin expression and fusion as seen in control cells (data not shown). This inhibition, up to 30%, was proportional to the increased expression of CDK4 in the transfected cells (Figure 6). Forced nuclear expression of CDK4 inhibits myogenic conversion of 10T1/2 cells in differentiation medium (DM) in the absence of cyclin Dl.

Therefore, the expression of CDK4 targeted to the nucleus, in the absence of cyclin Dl expression, can inhibit MyoD function in a concentration dependent manner, as measured by the reduction in the myogenic conversion of 10T1/2 mouse fibroblasts. From this result we conclude that activated CDK4 is not required to inhibit myogenesis since cyclin Dl is not expressed in cells maintained in differentiation medium. Although cyclin D3 expression is upregulated in differentiated muscle cultures, the cytoplasmic location of CDK4 in myotubes suggests that an active cyclin D3-CDK4 nuclear complex is not formed in low

mitogen medium. Therefore, we assume the forced nuclear localization of CDK4, in the absence of endogenous cyclin Dl, does not form an active kinase with cyclin D3.

## VIII. Phosphorylation

5

15

20

30

35

In this Example, the inventors examined whether an active CDK4 could phosphorylate MyoD directly to inhibit MyoD transactivation functions, by first examining whether or not MyoD could be phosphorylated by CDK4.

## Production of Activated CDK4

Activated CDK4 was prepared from baculovirus infected Sf9 cell extracts essentially as described in Example I.

## **Phosphorylation**

An <u>in vitro</u> kinase assay was performed using activated, immunoprecipitated CDK4 prepared from Sf9 cell extracts. 1–5 μl of Sf9 cell extract was mixed with 10 μl protein–G slurry (1:1 by volume of protein G suspension) washed with kinase buffer (50 mM Hepes, pH 7.5, 10mM MgCl2, 1mM DTT) and l μl of antibody (Santa Cruz; cat. no. sc–260–G) for one hour on ice. The slurry was then washed three times with 1 ml kinase buffer and then mixed with the l00 ng of either GST–RB target (pRB, amino acids 769–928, Kato et al., <u>Genes Dev.</u>, 1993, 7(3): p. 331–342) or renatured his–tagged MyoD plus or minus 200 ng of GST–pl6 (Serrano et al., <u>Nature</u>, 1993, **336**(6456): p.704–707) incubated in 20 μl of kinase buffer and assayed as described by Kato et al., <u>supra</u>.

## 25 Results

RB phosphorylation was sensitive to the CDK4 inhibitor, pl6 (Figure 7, lanes 1 to 3) whereas MyoD phosphorylation was essentially insensitive to pl6 inhibition (Figure 7, lanes 4 and 5) and was considered CDK4-independent. 2D-peptide maps for MyoD, +/- p16, using Glu-C and proteinase K reveal no qualitative differences in the phosphopeptide pattern (data not shown). Phosphorylation inhibited by the addition of pl6 to the kinase reaction, a specific CDK4 inhibitor, was assumed to be the CDK4-dependent component of the phosphorylation reaction.

Of the RB phosphorylation observed, 90–95% was inhibited by pl6 (Figure 7, lanes 2 and 3) whereas only 10–20% of the MyoD phosphorylation was sensitive to pl6 in the reaction, as determined by phosphoimager analysis (Figure 7, lanes 4

and 5). Two-dimensional phosphopeptide maps of MyoD protein, phosphorylated in the presence and absence of pl6 and digested either with Glu-C or proteinase K, also demonstrated no qualitative differences in the phosphopeptide patterns for either enzyme (data not shown). We assume that the non-CDK4-type kinases coprecipitated nonspecifically with the CDK4 antibody are responsible for the pl6-insensitive MyoD phosphorylation since both pl6 and MyoD bound to CDK4 inhibit kinase activity. Our data do not support the hypothesis that cyclin Dl inhibition of myogenesis is associated with a CDK4-dependent phosphorylation of MyoD since CDK4, in the absence of cyclin Dl, can block both MyoD DNA-binding and the myogenic conversion of 1OT1/2 cells, and MyoD is a very poor substrate for CDK4 phosphorylation in vitro compared to RB.

## IX. Mammalian Two-Hybrid System

To determine whether the MyoD-RB reaction is dependent upon the expression levels of MyoD in the cell, the mammalian two-hybrid system was used to analyze MyoD, myogenin and RB interactions <u>in vivo</u> in C2C12 muscle cells under growth and differentiation conditions.

## **Vector Preparation**

Chicken MyoD (CMD1) and myogenin were PCR amplified from pRSET clones (described by Shirakata, et al., Genes Dev, (1993), 7(12A): 2456–70) using Pfu polymerase (Stratagene), a 5' primer containing a BamH1 site and a 3' primer with a Hind3 site, and cloned into the corresponding sites in the vp16 vector pVP16 (Clonetech). E2F was cut from CMV–E2F (described by Helin, et al., (1993), Mol Cell Biol, 13(10): 6501–8) with BamH1, flush ended with T4 DNA polymerase and cloned into pVP16 that had been cut with Mlul and flush ended. Human RB (described by Kaye, et al., (1990), Proc Natl Acad Sci USA, 87(17): 6922–6) was cloned as a flush BamH1/Asp718 fragment into the flush Mlul site in the gal4 vector pM (Clonetech). Chicken E12 was cloned into the gal4 DNA–binding domain vector, pM. The sequence of chicken E12 is unpublished (Paterson lab). However, it is very similar to the human sequence, which is published, and the human sequence could be used in its place. Myogenin (ch—myogenin), (gene bank accession number D90157) was inserted into the vp16 activation domain plasmid, pVP16.

#### Cell Culture and Transfection

C2C12 myoblasts (3x10<sup>5</sup> cells per 60 mm dish or multiwell plate) were grown in DMEM with 20% fetal calf serum and transfected with the vectors described above using Fugene–6 (Boehringer Mannheim) according to the manufacturers directions.

1  $\mu$ g of gal4 luciferase reporter, 2  $\mu$ g each of the gal4 and vp16 constructs, and 300 ng of beta galactosidase expression plasmid, pCH110 (Pharmacia), for a total of 5.3  $\mu$ g DNA, was used for each transfection.

## 10 <u>Cell Harvest and Assay</u>

5

15

20

25

30

35

24-36 hours after transfection, the cells were placed in 2% horse serum plus insulin (10 µg/ml) to induce differentiation. After an additional 24–48 hours, cells were harvested for the luciferase reporter assay. Luciferase assays were performed using a Promega kit (cat. no. E4030) following the manufacturer's instructions and analyzed on a Victor 1420 multi label counter.

#### Results

Even in the presence of measurable endogenous levels of RB and E2F, both RB and E2F demonstrated significant interaction in C2C12 cells under growth conditions (10–15 fold activation of the luciferase reporter). Additionally, these proteins demonstrated significant interaction in low serum medium (used to induce muscle cell differentiation) See Figure 8a, line 7.

In contrast, there was no evidence for any interaction between RB and either MyoD or myogenin (ch-myogenin) in high or low serum culture medium (Figure 8a, lanes 3 and 5). However, all the control interactions were clearly seen: Both the vp16 MyoD(CMD1) and vp16 myogenin(ch-myogenin) constructs interacted strongly with gal4 E12 (Figure 8b, lanes 4 and 5), a known partner for these myogenic factors. In addition, both the MyoD and myogenin vp16 constructs were capable of efficiently converting 10T1/2 mouse fibroblasts to muscle (data not shown).

Despite previous reports, these results do not support the hypothesis that either MyoD or myogenin interact with RB to regulate cell cycle decisions in the terminal differentiation of the myoblast. C2C12 cells can differentiate normally under conditions that provide no evidence for an interaction between the myogenic factors and RB.

25

30

## XI. Mapping the MyoD CDK4 binding domain

Preliminary results demonstrated that MyoD binds specifically to CDK4 expressed either in baculovirus infected Sf9 cells or in bacteria, either as a CDK4/cyclin D1 complex or CDK4 alone (data not shown). Thus, the potential MyoD CDK4-binding domain was mapped.

## Preparation of MyoD-GST fusion proteins

Various domains of MyoD(CMD1), shown in Figure 9 (NH<sub>2</sub>, bHLH and

COOH fragments) and indicated by amino acid number from chicken MyoD, were amplified by PCR with pPfu with 5' primers containing a BamH1 site and 3' primers with an EcoR1 site and were cloned into the same sites either in pCDNA3

(Invitrogen) or GST-pCEFL (a eukaryotic GST expression vector from S. Gutkind). An NLS (MCPKKRKV) was incorporated into the 5' primer when the amplified fragment did not have a nuclear localization signal.

## **Fusion Protein Expression**

Proteins were expressed in E. coli strain BL21 (DE3) and purified as described by Shirakata et al., <u>Genes Dev</u>. 1993 7(12A):2456-70. The MyoD -GST fusion proteins were bound to glutathione agaraose.

#### **Binding Experiments**

The binding experiments were performed with CDK4 produced in the baculovirus system (Kato et al., <u>Genes Dev.</u> 1993 7(3):331–42) essentially as described by Kaelin et al., <u>Cell</u> 1991 64(3):521–32.

## **Deletion mapping**

Deletion mapping of the MyoD (CMD1) carboxy terminus narrowed the CDK4-binding region to a maximum of fifteen amino acids between residues 189–203, well outside the bHLH domain (Figure 10b, lanes 4 and 5). To confirm that these fifteen amino acids bound CDK4, the fifteen amino acid peptide was joined to glutathione transferase and tested for its ability to bind the CDK4 from Sf9 cell extracts.

# Results

Unexpectedly, CDK4 bound to the carboxy terminus of MyoD (CMD1) in the absence of the bHLH domain (Figure 10a, lane 5). The fifteen amino acid domain in the carboxy terminus of MyoD (CMD1) bound CDK4 as efficiently as full-sized MyoD (CMD1) (Figure 10c, lanes 2 and 3).

# XII. MyoD inhibition of RB

To determine whether MyoD could inhibit CDK4 activity towards RB in addition to the ability of CDK4 to influence MyoD function, MyoD(CMD1) was assayed in a CDK4-dependent phosphorylation reaction for its ability to inhibit the phosphorylation of GST-RB in vitro.

#### Transfection and Cell culture

Sf9 cells were infected with wild-type virus, cyclin D1 or CDK4 encoding virus stocks. After incubation for 48 hours at 23° C, cell extracts were made using methods known to those of skill in the art.

# Phosphorylation assay

Extracts produced from Sf9 cells coinfected with virus stocks encoding cyclin D1 and CDK4 were used as a source of activated kinase. Antibody to phosphoserine 780, a CDK4-specific site in RB, was used to measure phosphorylation. Myogenin was used as a control since it does not bind MyoD (Figure 10a, lane 3). Knudsen, et al., *Cell Biol*, (1997),17(10):5771-83; and Kitagawa, et al., *Embo J*, (1996),15(24):7060-9.

25

5

10

15

#### Results

Extracts from Sf9 cells coinfected with wild-type virus (Figure 11a, lane 1), a cyclin D1 encoding virus (Figure 11a, lane 2) or a CDK4 encoding virus (Figure 11, lane 3) give background levels of kinase activity on serine 780 (antibody to phosphoserine 780). Coinfections with baculovirus stocks encoding CDK4 and cyclin D1 produce active CDK4 (Figure 11a, lane 4) that specifically phosphorylates RB (GST-RB (aa 767-928)) on serine 780. Phosphorylation is inhibited to background levels (e.g., by more than 90%) by the CDK4-specific kinase inhibitor, p16 (Figure 11a, lane 5, added as GST-p16).

30

# XIII. Demonstration of CDK4 Binding Peptide Inhibitory Properties

In this experiment, both the full-sized and the carboxy terminal domain of MyoD (CMD1), were titrated in the CDK4-dependent phosphorylation reaction to test their ability to inhibit kinase activity. Myogenin, which does not bind to CDK4, was used as the control. As before, CDK4-dependent phosphorylation of RB was measured using an antibody to phosphoserine 780.

As used herein, the term "amino terminus of MyoD" refers to amino acids 1–92 of CMD1. The "bHLH domain" refers to amino acids 93–153. The "carboxy terminus" refers to amino acids 153–298. The "fifteen amino acid CDK4-binding peptide" refers to amino acids 189–203. Amino acid 299 is a terminator codon.

# Transfection and Cell Culture

Active CDK4 was produced in Sf9 cells coinfected with baculovirus stocks encoding CDK4 and cyclin D1 (from C. J. Sherr).

15

20

25

30

35

10

5

# A. CDK4 Kinase Assay

The kinase assay was performed essentially as described by Kato, et al., *Genes Dev*, (1993), 7(3):331–42. The CDK4–specific kinase inhibitor p16 (from D. Beach, [as described in Serrano, et al., *Nature*, (1993), 366(6456): 704–7]) was used as a GST–p16 fusion added to the kinase reaction. The GST–RB fusion (767–928) was used as a kinase target as previously described Kato, et al., *Genes Dev*, (1993), 7(3):331–42.

To test for kinase inhibition, the various proteins and protein domains were added to the reaction in two-fold increments from 75 ng to 300ng. For the pCDNA3 constructs, total protein concentration was kept constant in the kinase reaction by adding a 6-his tagged amino terminus of MyoD (CMD1) (aa 1-92), which does not bind CDK4. For the GST constructs, total protein concentration was kept constant in the kinase reaction by adding GST protein. After the assay the total reaction was run on SDS PAGE gels and analyzed by Western blot analysis. Specific phosphorylation of RB was measured with rabbit polyclonal antibodies to

Results

As shown in Figure 11b, full-length MyoD(CMD1) (second row), the C-terminus of MyoD(CMD1) (third row) and the fifteen amino acid CDK4-binding

RB phosphoserines 780 or 795 (New England Biolabs).

peptide fused to GST (fourth row) inhibit the CDK4-dependent phosphorylation of RB(GST-RB) in vitro. Myogenin (first row) (ch-myogenin), which does not bind to CDK4, had no inhibitory effect (Figure 11b, top row). Similarly, neither the N-terminus (amino acids 1–92) nor the bHLH domain of MyoD(CMD1) inhibited CDK4 activity (data not shown). GST protein alone was not inhibitory and behaved similar to myogenin (data not shown). Identical results were obtained using an antibody to phosphoserine 795, another CDK4 dependent phosphorylated residue in RB (data not shown).

Therefore, the observed MyoD-CDK4 interaction <u>in vitro</u> can specifically inhibit the phosphorylation of RB in a dose-dependent manner on serine residues known to be targeted by CDK4. Thus, forced expression of either MyoD or its CDK4-binding subdomain should inhibit CDK4 in cells to block the phosphorylation of RB and arrest growth, similar to the growth inhibitory effects of full-length MyoD.

15

20

25

10

5

# B. BrdU incorporation assay

A BrdU incorporation assay was employed to determine whether expression of either the carboxy terminus or the minimal fifteen amino acid CDK4-binding domain of MyoD(CMD1) could induce growth arrest in cultured cells.

# Transfection & Cell Culture

10T1/2 cells in growth medium were cotransfected with plasmids expressing full-sized MyoD(CMD1) or the indicated domains, either as GST fusions in GST-pCEFL or as subclones in pCDNA3, along with a lacZ expression construct, pCH110, to mark transfected cells. The various domains were targeted to the nucleus with a nuclear tag in the absence of an existing NLS.

# BrdU assay

Cultures were maintained in 20% fetal calf serum. 72 hours after transfection cultures were refed with the same medium and 12 hours later they were given a one hour pulse with BrdU (10μM) prior to fixation, as previously described by Crescenzi et al., PNAS USA 1990 87(21):8442–6; and Sorrentino et al., Nature, 1990 345(6278):813–5. Cells were then reacted with rabbit anti–BrdU antibody (Jackson labs) and Xgal.

15

20

# Results

A typical result used in the scoring assay (Sorrentino et al., <u>Nature</u>, 1990, **345**(6278): p. 261–72 and Crescenzi et al., <u>Proc. Natl. Acad. Sci. USA</u>, 1990,

- 87(21): p. 8442-6) to measure the percentage of BrdU positive cells with the transfected cells expressing the various domains of MyoD labeled cells (indicating the cotransfected beta galactosidase expression vector) is shown. All the GST-MyoD domain constructs used were localized to the nuclei of transfected cells, as typified for the fifteen amino acid CDK4-binding peptide fusion with an NLS.
- Neither the empty vectors (pCDNA3 or GST-pCEFL with an NLS) nor the amino terminal portion of MyoD(CMD1) (amino acids 1-92 in pCDNA3 with an NLS) were able to inhibit cell growth (Figure 12). In contrast, full MyoD(CMD1) showed growth inhibitory activity (Figure 12).

The amino terminus of MyoD(CMD1) (amino acids 1–92) was ineffective in growth inhibition whereas the bHLH domain (amino acids 93–153) and, to an even greater extent, the carboxy terminus (amino acids 153–298) were both effective at inhibiting cell growth (Figure 12, upper section). Again, the fifteen amino acid CDK4–binding peptide from the carboxy terminus (amino acids 189–203) was as potent as full–sized MyoD(CMD1) in growth inhibitory activity in this assay (Figure 12, lower section). The bHLH domain of MyoD alone also triggered growth arrest, although to a lesser extent.

# C. Phosphorylation of RB

This experiment is designed to confirm that growth arrest correlates with a change in the phosphorylation status of RB induced by MyoD.

# Vector preparation

The inhibitory activity of the fifteen amino acid CDK4 binding peptide on RB phosphorylation in cells was examined by fusing the fifteen amino acid CDK4–30 binding peptide of MyoD with a nuclear localization tag to the carboxy terminus of GFP and used in the same transfection protocol with GST–RB.

# Transfection and Cell Culture

10T1/2 cells were cotransfected with a GST-RB target (amino acids 767–928) containing a nuclear localization tag, and either MyoD(CMD1) or the carboxy terminus of MyoD with an NLS.

5

# Recovery of GST-RB

48 hours after transfection, nuclear extracts of the 10T1/2 cells were prepared. The nuclear extracts were passed over glutathione agaraose to capture the GST-RB target.

10

15

25

# Western Blotting

Western blot analysis of the captured GST-RB target was performed with phosphoserine specific antibodies to the CDK4 target sites, serines 780 and 795. Gel loadings for the phosphoserine antibody analysis were adjusted to the same amount of GST-RB target protein by preliminary western analysis of the extracts (Figure 13).

# Results

Both the full sized and the carboxy terminal domain of MyoD (CMD1)

reduced the phosphorylation of serines 780 and 795 on the cotransfected RB target protein, as shown for the C-terminus with antibodies to phosphoserines 780 and 795 (Figure 13)

# D. Growth Inhibition

This experiment was designed to test the inhibitory activity of the fifteen amino acid CDK4-binding peptide on RB phosphorylation in cells.

# Transfection

C2C12 myoblasts were cotransfected with a GST-pCEFL RB expression plasmid (RB aa 767-928) and pCDNA3 expression plasmids for full-sized or the C-terminus of MyoD(CMD1). GST-RB was recovered from the cells using methods known to those of skill in the art. The samples were adjusted for equal levels GST protein and loaded on SDS-PAGE gels for western analysis by phosphoserine 780 and 795-specific antibodies.

# Results

The 15 amino acid domain in the carboxy terminus of MyoD that binds CDK4 also inhibits CDK4 kinase activity and cell growth when expressed in cell nuclei.

The entire disclosures of all citations cited throughout the specification, and the references cited therein, are hereby expressly incorporated by reference.

Although the foregoing refers to particular preferred embodiments, it will be understood that the present invention is not so limited. It will occur to those ordinarily skilled in the art that various modifications may be made to the disclosed embodiments without diverting from the overall concept of the invention. All such modifications are intended to be within the scope of the present invention. Thus only such limitations as appear in the appended claims should be placed upon the scope of the present invention.

15

10

5

# WHAT IS CLAIMED IS:

- 1. An isolated CDK4 binding peptide comprising:
- Tyr-Ser-Gly-Pro-Pro-Xaa<sub>1</sub>-Xaa<sub>2</sub>-Xaa<sub>3</sub>-Arg-Arg-Xaa<sub>4</sub>-Asn-Xaa<sub>5</sub>-Tyr-Xaa<sub>6</sub> wherein Xaa<sub>1</sub>= Cys or Ser; Xaa<sub>2</sub>= Ser or Gly; Xaa<sub>3</sub>= Ser, Ala or Pro; Xaa<sub>4</sub>=Arg or Gln; Xaa<sub>5</sub>= Ser, Cys or Gly; and Xaa<sub>6</sub>=Asp or Glu.
- 2. The isolated CDK4 binding peptide, wherein the isolated peptide does not include a bHLH domain.
- 3. A fusion protein comprising:
  - (a) a CDK4 binding peptide comprising Tyr-Ser-Gly-Pro-Pro-Xaa<sub>1</sub>-Xaa<sub>2</sub>-Xaa<sub>3</sub>-Arg-Arg-Xaa<sub>4</sub>-Asn-Xaa<sub>5</sub>-Tyr-Xaa<sub>6</sub> wherein Xaa<sub>1</sub>= Cys or Ser; Xaa<sub>2</sub>= Ser or Gly; Xaa<sub>3</sub>= Ser, Ala or Pro; Xaa<sub>4</sub>=Arg or Gln; Xaa<sub>5</sub>= Ser, Cys or Gly; and Xaa<sub>6</sub>=Asp or Glu; and
  - (b) a heterologous amino acid sequence.
- 4. The fusion protein of claim 3 wherein the heterologous amino acid sequence comprises a nuclear localization signal.
- 5. The fusion protein of claim 4 wherein the CDK4 binding peptide is at the C-terminus of the fusion protein.
- 6. A nucleotide sequence encoding a CDK4 binding peptide, wherein the CDK4 binding peptide comprises:

Tyr-Ser-Gly-Pro-Pro-Xaa<sub>1</sub>-Xaa<sub>2</sub>-Xaa<sub>3</sub>-Arg-Arg-Xaa<sub>4</sub>-Asn-Xaa<sub>5</sub>-Tyr-Xaa<sub>6</sub> wherein Xaa<sub>1</sub>= Cys or Ser; Xaa<sub>2</sub>= Ser or Gly; Xaa<sub>3</sub>= Ser, Ala or Pro; Xaa<sub>4</sub>=Arg or Gln; Xaa<sub>5</sub>= Ser, Cys or Gly; and Xaa<sub>6</sub>=Asp or Glu.

- 7. The nucleotide sequence of claim 6, wherein the nucleotide sequence is a DNA sequence.
- 8. A method of inhibiting cell growth, comprising:
  administering to a patient a CDK4 binding peptide comprising:

  Tyr-Ser-Gly-Pro-Pro-Xaa<sub>1</sub>-Xaa<sub>2</sub>-Xaa<sub>3</sub>-Arg-Arg-Xaa<sub>4</sub>-Asn-Xaa<sub>5</sub>-Tyr-Xaa<sub>6</sub>

wherein Xaa<sub>1</sub>= Cys or Ser; Xaa<sub>2</sub>= Ser or Gly; Xaa<sub>3</sub>= Ser, Ala or Pro; Xaa<sub>4</sub>=Arg or Gln; Xaa<sub>5</sub>= Ser, Cys or Gly; and Xaa<sub>6</sub>=Asp or Glu in an amount effective to inhibit cell growth.

9. A method of inhibiting the activity of CDK4 comprising: contacting CDK4 with a CDK4 binding peptide comprising:

Tyr-Ser-Gly-Pro-Pro-Xaa<sub>1</sub>-Xaa<sub>2</sub>-Xaa<sub>3</sub>-Arg-Arg-Xaa<sub>4</sub>-Asn-Xaa<sub>5</sub>-Tyr-Xaa<sub>6</sub> wherein Xaa<sub>1</sub>= Cys or Ser; Xaa<sub>2</sub>= Ser or Gly; Xaa<sub>3</sub>= Ser, Ala or Pro; Xaa<sub>4</sub>=Arg or Gln; Xaa<sub>5</sub>= Ser, Cys or Gly; and Xaa<sub>6</sub>=Asp or Glu in an amount effective to inhibit the activity of CDK4.

5

# 42 ABSTRACT

The provides a CDK4 binding peptide, and a nucleic acid sequence coding therefore, that is capable of specifically binding cyclin dependent kinase (CDK4) to inhibit CDK4 activity and cell growth. The invention also includes variants of the CDK4 binding peptide which comprise polypeptides which have at least about 80% amino acid sequence identity with the amino acid sequence of the CDK4 binding peptide. In another embodiment, the invention provides chimeric molecules comprising a CDK4 binding peptide fused to a heterologous peptide or amino acid sequence, preferably a nuclear localization signal. Therapeutic and diagnostic methods are also provided.

Sheet 1 of 14

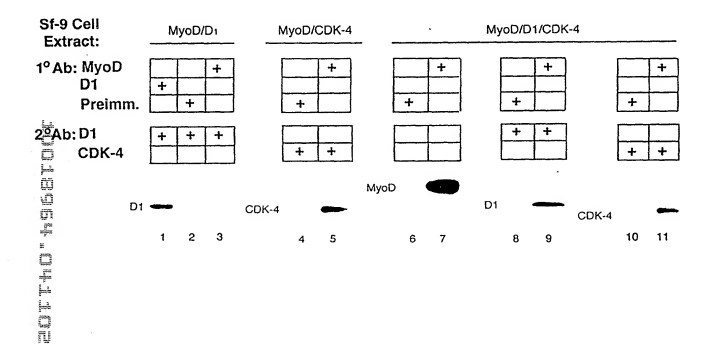


FIG. 1

Inventor: PATERSON et al. Docket No.: 11613.33USWO Title: CDK4 BINDING PEPTIDE

Attorney Name: Melissa J. Pytel (Reg. No. 41,512) Phone No: 612.371.5304 Sheet 2 of 14

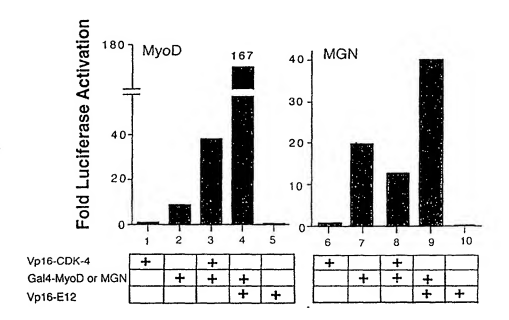


FIG. 2

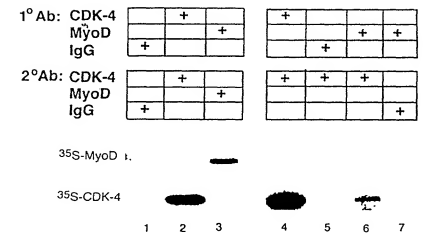


FIG. 3

Sheet 4 of 14

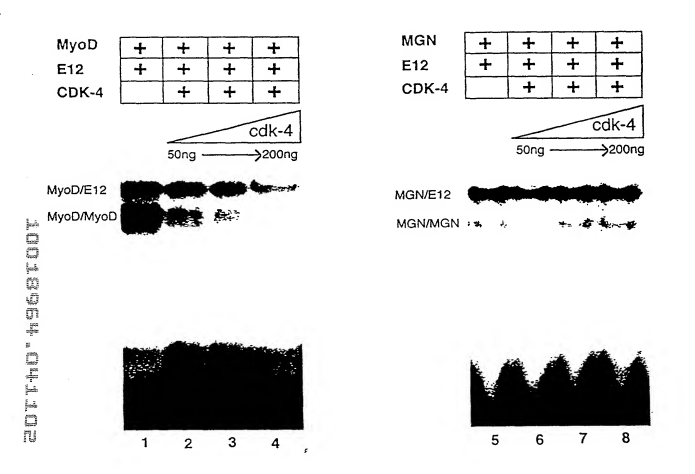
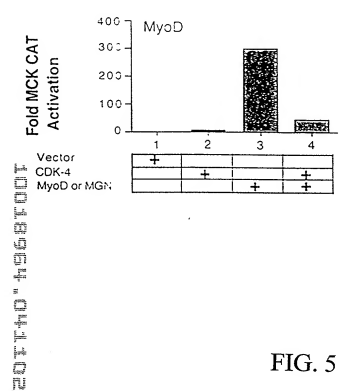


FIG. 4

Inventor: PATERSON et al.
Docket No.: 11613.33USWO
Title: CDK4 BINDING PEPTIDE
Attorney Name: Melissa J. Pytel (Reg. No. 41,512)
Phone No.: 612.371.5304
Sheet 5 of 14

018964 10





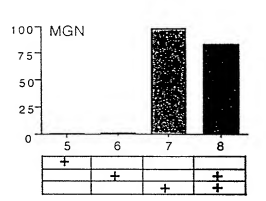


FIG. 5

Inventor: PATERSON et al.
Docket No.: 11613.33USWO
Title: CDK4 BINDING PEPTIDE
Attorney Name: Melissa J. Pytel (Reg. No. 41,512)
Phone No.: 612.371.5304
Sheet 6 of 14

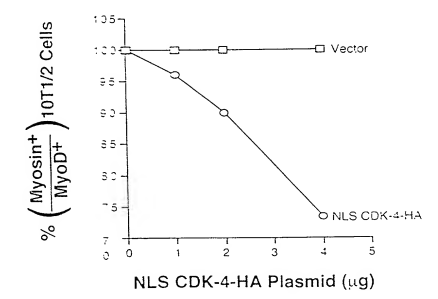
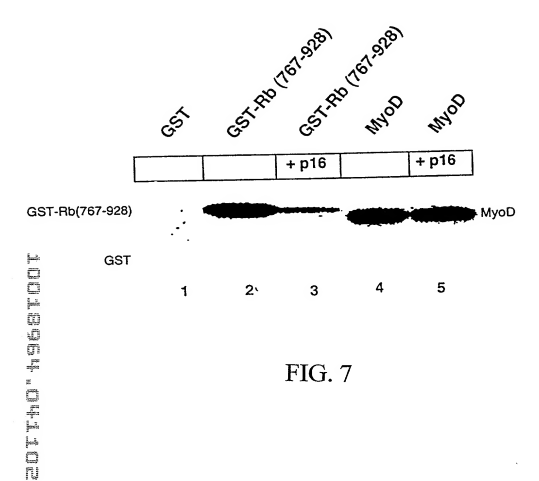


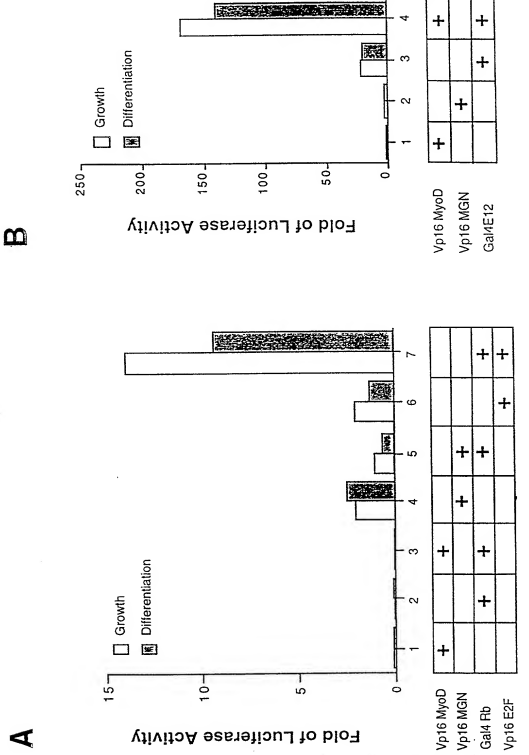
FIG. 6



496810/01

Inventor: PATERSON et al.

Docket No.: 11613-33USWO
Title: CDK4 EINDING PEPTIDE
After No.: 612.371.5304
Phone No.: 612.371.5304
Sheet 8 of 14



+

FIG. 8

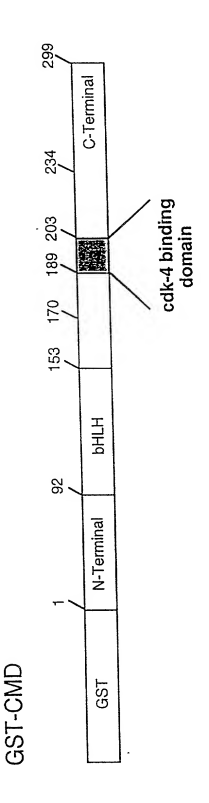
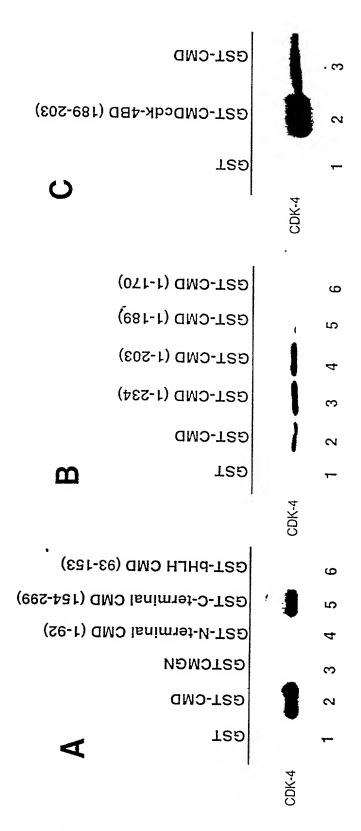


FIG. 9

Phone No.: 612.371.5304

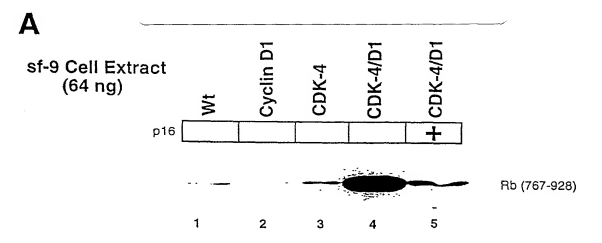
Inventor: PATERSON et al.

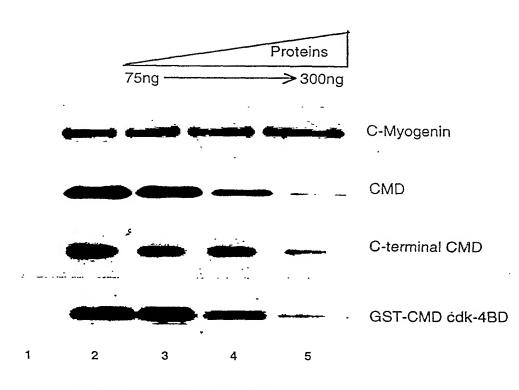
Title: CDK4 BINDING PEPTIDE
Attorney Vanne: Melissa J. Pyrel (Reg. No. 41,512)



IG. 10

B





PO4-Rb(Anti-P780)

FIG. 11

%Inhibition	% 0	28 %	% 7	34 %	47 %	% 0	41 %	
wl%		u,						
No. BrdU+ Cells 500 β-gal+ Cells (n=6)	88.4 ± 6.2	$37.1 \pm 4.2$	86.8 ± 4.8	58.4 ± 3.9	<b>46.9</b> ± <b>6.4</b>	96.7 ± 5.9	57.4 ± 4.3	
CMD aa	Vector	с-смр 1-299	1-92	93-153	153-299	1-92	189-203	
		N-CMD BHLH	N-CMD	РНГН	C-CMD	GST. N-CMD	SC TSD	
CDK-4 Binding		+		1	+	1	+	

FIG. 12

Sheet 12 of 14

Title: CDK4 BINDING PEPTIDE

Title: To f 14

Inventor: PATERSON et al.
Docket No.: 11613.33USWO
Title: CDK4 BINDING PEPTIDE
Attorney Name: Melissa J, Pytel (Reg. No. 41,512)
Phone No.: 612.371.5304
Sheet 13 of 14

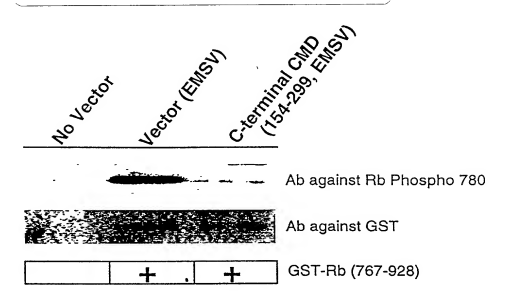


FIG. 13

Inventor: PATERSON et al.
Docket No.: 11613.33USWO
Title: CDK4 BINDING PEPTIDE
Attorney Name: Melissa J. Pytel (Reg. No. 41,512)
Phone No.: 612.371.5304

Sheet 14 of 14

**SPECIES** 

PEPTIDE SEQUENCE

Chicken Human Rat

Murine

Pig

**YSGPPCSSRRRNSYDS YSGPPSGARRRNCYEG** YSGPPSGPRRQNGYDA **YSGPPSGPRRQNGYDT YSGPPSGARRRNCYDG** 

Consensus

YSGPPC/SG/S-RRR/QN-YD/E-

Xenopus-A Xenopus-B Zebra fish

YNSPPCGSRRRNSYD YNSPPCSSRRRNSYD **FMGPTCQTRRRNSYD** 

FIG. 14

#### **United States Patent Application**

#### COMBINED DECLARATION AND POWER OF ATTORNEY

As a below named inventor I hereby declare that: my residence, post office address and citizenship are as stated below next to my name;

that			
I verily believe I am the original, the below) of the subject matter which	first and sole inventor (if only on a is claimed and for which a pate	ne name is listed below) or a joint in ent is sought on the invention entitle	ventor (if plural inventors are named d: CDK4 BINDING PEPTIDE
The specification of which			
a.  is attached hereto			
PCT-filed application) de		ial no. 10/018964 and was amended ional no. PCT/US00/14489 filed 25 I ited States patent.	
I hereby state that I have reviewed any amendment referred to above.		the above-identified specification, in	ncluding the claims, as amended by
I acknowledge the duty to disclose Federal Regulations, § 1.56 (attack		to the patentability of this application	n in accordance with Title 37, Code of
	lso identified below any foreign		pplication(s) for patent or inventor's certificate having a filing date before
a. no such application	ns have been filed.		
a. ☐ no such application  b. ☐ such applications	have been filed as follows:		
FOR	REIGN APPLICATION(S), IF ANY, (	CLAIMING PRIORITY UNDER 35 USC	§ 119
COUNTRY	APPLICATION NUMBER	DATE OF FILING (day, month, year)	DATE OF ISSUE (day, month, year)
ALL FOR	 FIGN APPLICATION(S), IF ANY, F	ILED BEFORE THE PRIORITY APPLIC	CATION(S)
COUNTRY	APPLICATION NUMBER	DATE OF FILING (day, month, year)	DATE OF ISSUE (day, month, year)
			1

I hereby claim the benefit under Title 35, United States Code, § 120/365 of any United States and PCT international application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35, United States Code, § 112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, § 1.56(a) which occurred between the filing date of the prior application and the national or PCT international filing date of this application.

U.S. APPLICATION NUMBER	DATE OF FILING (day, month, year)	STATUS (patented, pending, abandoned)

I hereby claim the benefit under Title 35, United States Code § 119(e) of any United States provisional application(s) listed below:

U.S. PROVISIONAL APPLICATION NUMBER	DATE OF FILING (Day, Month, Year)
60/139934	18 June 1999

Power of Attorney: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith:

James C. Haight, 25,588; Robert Benson, 33,612; Jack Spiegel, 34,477; Susan S. Rucker, 35,762; David R. Sadowski, 32,808; Steven M. Ferguson, 38,448; John Peter Kim, 38,514; Stephen L. Finley, 36,357; Norbert Pontzer, 40,777; Richard Rodriquez, 45,980; and Marlene Shinn, 46,005.

all of the Office of Technology Transfer, National Institutes of Health, Rockville, Maryland; with an Associate Power of Attorney to:

Brian H. Batzli, 32,960; Steven C. Bruess, 34,130; James R. Chiapetta, 39,634; Ronald A. Daignault, 25,968; Dennis R. Daley, 34,994; Mark J. DiPietro, 28,707; Alan G. Gorman, 38,472; Richard J. Gregson, 41,804; John J. Gresens, 33,112; Randall A. Hillson, 31,838; Denise M. Kettelberger, 33,924; Alan W. Kowalchyk, 31,535; Katherine M. Kowalchyk, 36,848; Andrew J. Leon, 46,869; Douglas P. Mueller, 30,300; Bryan K. Phillips, 46,990; Paul J. Prendergast, 46,068; Melissa J. Pytel, 41,512; John C. Reich, 37,703; Ted R. Rittmaster, 32,933; Kirstin L. Stoll-DeBell, 43,164; John P. Sumner, 29,114; David K. Tellekson, 32,314; Karrie Weaver, 43,245; Brian Whipps, 43,261; and J. Scot Wickhem, 41,376.

SEND CORRESPOND	ENCE TO:		DIRECT TELEPHONE CALLS TO:			
Merchant & Gould P.C.			Melissa J. Pytel			
3200 IDS Center			(612) 371.5304			
80 South Eighth S	The company of the Co					
Minneapolis, MN	\$5402-2215					
Full Name	Family Name	First	t Given Name	1	Second Given Name	
2 Of Inventor	PATERSON	Bruc	e		M.	
0 Residence	City	State	e or Foreign Country		Country of Citizenship	
La Citizenship	Bethesda	Mary	yland (V)		United States	
1 Mailing	Address	City			State & Zip Code/Country	
Address	9806 Holmhurst Road	Beth	esda		Maryland 20817 / United States	
Full Name	Family Name	First	t Given Name		Second Given Name	
.2 Of Inventor	ZHANG.	Jian-	Min			
			<u> </u>			
0 Residence	City	State	e or Foreign Country		Country of Citizenship	
& Citizenship	Richmond	Virg	inia //		China	
2 Mailing	Address	City			State & Zip Code/Country	
Address	900 Pump Road, #4		mond		Virginia 23233 / United States	
I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and believe						
are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so						
made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful						
false statements may jeopardize the validity of the application or any patent issued thereon.						
Signature of Inventor 201: Dates Parch 25, 2502						
Signature of Inventor 202:				Date:		

Power of Attorney: As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith:

James C. Haight, 25,588; Robert Benson, 33,612; Jack Spiegel, 34,477; Susan S. Rucker, 35,762; David R. Sadowski, 32,808; Steven M. Ferguson, 38,448; John Peter Kim, 38,514; Stephen L. Finley, 36,357; Norbert Pontzer, 40,777; Richard Rodriquez, 45,980; and Marlene Shinn, 46,005.

all of the Office of Technology Transfer, National Institutes of Health, Rockville, Maryland; with an Associate Power of Attorney to:

Brian H. Batzli, 32,960; Steven C. Bruess, 34,130; James R. Chiapetta, 39,634; Ronald A. Daignault, 25,968; Dennis R. Daley, 34,994; Mark J. DiPietro, 28,707; Alan G. Gorman, 38,472; Richard J. Gregson, 41,804; John J. Gresens, 33,112; Randall A. Hillson, 31,838; Denise M. Kettelberger, 33,924; Alan W. Kowalchyk, 31,535; Katherine M. Kowalchyk, 36,848; Andrew J. Leon, 46,869; Douglas P. Mueller, 30,300; Bryan K. Phillips, 46,990; Paul J. Prendergast, 46,068; Melissa J. Pytel, 41,512; John C. Reich, 37,703; Ted R. Rittmaster, 32,933; Kirstin L. Stoll-DeBell, 43,164; John P. Sumner, 29,114; David K. Tellekson, 32,314; Karrie Weaver, 43,245; Brian Whipps, 43,261; and J. Scot Wickhem, 41,376.

SEND CORRESPONDENCE TO:			DIRECT TELEPHONE CALLS TO:				
Merchant & Gould P.C.				Melissa J. Pytel			
3200 IDS Center				(612) 371.5304			
JI *	80 South Eighth Street						
Mi	ineapolis, MN	55402-2215					
417	Full Name	Family Name	Firs	t Given Name		Second Given Name	
2	Of Inventor	PATERSON	Bruc	ce		M.	
1	*			***			
0 .	Residence	City	Stat	e or Foreign Country		Country of Citizenship	
	& Citizenship	Bethesda	Mar	yland		United States	
1 1	Mailing	Address Cit		City		State & Zip Code/Country	
	Address	9806 Holmhurst Road	Beth	nesda		Maryland 20817 / United States	
Ē	Full Name	Family Name	Firs	t Given Name		Second Given Name	
2 1	Of Inventor	ZHANG		Jian-Min			
u will	#						
0 🗓	. Residence	City	Stat	e or Foreign Country		Country of Citizenship	
	& Citizenship	Richmond	Virg	inia		China	
2	Mailing	Address	City	,		State & Zip Code/Country	
	Address	900 Pump Road, #4		mond		Virginia 23233 / United States	
	I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belie						
	are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so						
	made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful						
false	false statements may jeopardize the validity of the application or any patent issued thereon.						
Signature of Inventor 201: Date:							
Sign	ature of Inventor 2	02: 3		Date: -2	123/02		
<u>.</u>		= 5- 500 0		7	122100		
	· · · · · · · · · · · · · · · · · · ·						

#### SEQUENCE LISTING

```
<110> Paterson, Bruce M.
           Zhang, Jian-Min
    <120> CDK4 BINDING PEPTIDE
    <130> 11613.33USWO
    <140> US 10/018,964
    <141>
           2001-12-18
           PCT/US00/14489
    <150>
           2000-05-25
    <151>
    <150> US 60/139,934
           1999-06-18
    <151>
    <160>
    <170> PatentIn version 3.1
    <210>
           7
    <211>
          15
    <212>
          PRT
<213> Artificial Sequence
    <220>
    <223> Synthetic
    <220>
    <221> MISC_FEATURE
    <222>
           (6)..(6)
    <223> Xaa is an amino acid having an uncharged polar side chain, including
           Cys or Ser
    <220>
    <221>
          MISC_FEATURE
    <222>
           (7)..(7)
    <223>
          Xaa is an amino acid having an uncharged side chain, including Ser or
           Gly
    <220>
    <221>
          MISC_FEATURE
    <222>
           (8)..(8)
          Xaa is an amino acid having an uncharged side chain, including Ser, Ala
    <223>
    <220>
    <221> MISC FEATURE
    <222>
          (11)..(11)
   <223> Xaa is an amino acid having a polar side chain, including Arg or
          Gln
   <220>
   <221> MISC FEATURE
                                          1
```

```
<223> Xaa is an amino acid having an uncharged side chain, including Ser, Cys
    <220>
    <221>
           MISC FEATURE
    <222>
           (15)...(15)
           Xaa is an amino acid having a charged polar side chain, including
           Asp or Glu
    <400> 1
    Tyr Ser Gly Pro Pro Xaa Xaa Xaa Arg Arg Xaa Asn Xaa Tyr Xaa
                                        10
    <210> 2
    <211> 16
    <212> PRT
    <213> Gallus gallus
    <400> 2
Tyr Ser Gly Pro Pro Cys Ser Ser Arg Arg Arg Asn Ser Tyr Asp Ser
    <210> 3
    <211> 16
    <212> PRT
    <213> Homo sapiens
    <400> 3
    Tyr Ser Gly Pro Pro Ser Gly Ala Arg Arg Arg Asn Cys Tyr Glu Gly
    <210>
          4
    <211>
           16
    <212>
           PRT
    <213> Rattus rattus
    <400> 4
    Tyr Ser Gly Pro Pro Ser Gly Pro Arg Gln Asn Gly Tyr Asp Ala
    <210> 5
    <211>
           16
    <212> PRT
    <213> Mus musculus
    <400> 5
```

<222> (13)..(13)

Tyr Ser Gly Pro Pro Ser Gly Pro Arg Arg Gln Asn Gly Tyr Asp Thr

```
<210> 6
<211> 16
<212> PRT
<213> Sus scrofa
<400> 6
Tyr Ser Gly Pro Pro Ser Gly Ala Arg Arg Arg Asn Cys Tyr Asp Gly
<210> 7
<211> 15
<212> PRT
<213> Xenopus laevis
<400> 7
Tyr Asn Ser Pro Pro Cys Gly Ser Arg Arg Arg Asn Ser Tyr Asp
<210> 8
<211> 15
<212> PRT
<213> Xenopus laevis
<400> 8
Tyr Asn Ser Pro Pro Cys Ser Ser Arg Arg Asn Ser Tyr Asp
<210> 9
<211> 15
<212> PRT
<213> Brachydanio rerio
<400> 9
Phe Met Gly Pro Thr Cys Gln Thr Arg Arg Arg Asn Ser Tyr Asp
```

# United States Patent & Trademark Office Office of Initial Patent Examination -- Scanning Division



Application deficience  Page(s)	of		were not presen
for scanning.		(Document title)	,
☐ Page(s)	of		were not presen
for scanning.		(Document title)	

Scanned copy is best available. Some drawings are too doork.